

(12) INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(19) World Intellectual Property Organization
International Bureau



(43) International Publication Date
7 March 2002 (07.03.2002)

PCT

(10) International Publication Number
WO 02/18404 A2

(51) International Patent Classification⁷: **C07H 19/00**

(21) International Application Number: **PCT/EP01/09633**

(22) International Filing Date: 21 August 2001 (21.08.2001)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
0021285.2 30 August 2000 (30.08.2000) GB
0026611.4 31 October 2000 (31.10.2000) GB

(71) Applicant: **F. HOFFMANN-LA ROCHE AG [CH/CH];**
124, Grenzacherstrasse, CH-4070 Basle (CH)..

(72) Inventors: **DEVOS, Rene; 4 Salmon Close, Welwyn Garden City, Hertfordshire AL7 1TR (GB). DYMOCK, Brian, William; 15 Vesta Avenue, St. Albans, Hertfordshire AL1 2PJ (GB). HOBBS, Christopher, John; 9 Magnolia Close, Hertford, Hertfordshire SG13 7UR (GB). JIANG, Wen-Rong; 20 Salmon Close, Welwyn Garden City, Hertfordshire AL7 1TR (GB). MARTIN, Joseph, Armstrong; 10 The Chownes, West Common, Harpenden, Herts AL5 2BN (GB). MERRITT, John, Herbert; 23 Bush Spring, Baldock, Hertfordshire SG7 6QT (GB). NAJERA, Isabel; 49 Salisbury Avenue, St. Albans, Hertfordshire AL1 4TZ (GB). SHIMMA, Nobuo;**

Higashikaigan-Minami 2-11-19, Chigasaki-shi, Kanagawa-ken 253-0054 (JP). TSUKUDA, Takuo; 540-22 Rensyoji, Odawara-shi, Kanagawa-ken 250-0865 (JP).

(74) Agent: **MEZGER, Wolfgang; 124 Grenzacherstrasse, CH-4070 Basle (CH).**

(81) Designated States (*national*): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.

(84) Designated States (*regional*): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

— without international search report and to be republished upon receipt of that report

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: **NUCLEOSIDE DERIVATIVES**

(57) Abstract: Use of compounds of formula I (I), wherein R¹ is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy, halogen, cyano, isocyano or azido; R² is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine; R³ is hydrogen; or R² and R³ together represent =CH₂; or R² and R³ represent fluorine; X is O, S or CH₂; a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and B signifies a purine base B1 which is connected through the 9-nitrogen of formula (B1), wherein R⁴ is hydrogen, hydroxyl, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen or SH; R⁵ is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH; R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen, SH or cyano; R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl; R⁹ is hydrogen, alkyl or aryl; or B signifies an oxidised purine base B2 which is connected through the 9-nitrogen of formula (B2), wherein R⁴, R⁵ and R⁶ are as defined above; or B signifies a purine base B3 which is connected through the 9-nitrogen of formula (B3), wherein R⁴ and R⁶ are as defined above; R¹⁰ is hydrogen, alkyl or aryl; Y is O, S or NR¹¹; R¹¹ is hydrogen, hydroxy, alkyl, OR⁹, heterocycl or NR⁷R⁸; R⁷, R⁸ and R⁹ are as defined above; or B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula (B4), wherein Z is O or S; R¹² is hydrogen, hydroxy, alkyl, alkoxy, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH; R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen; R⁷, R⁸ and R⁹ are as defined above; or B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula (B5), wherein Y, Z, R¹⁰ and R¹³ are as defined above for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment. The invention is concerned with novel and known purine and pyrimidine nucleoside derivatives, their use as inhibitors of subgenomic Hepatitis C Virus (HCV) RNA replication and pharmaceutical compositions of such compounds.

WO 02/18404 A2

Nucleoside Derivatives

The invention relates to nucleoside derivatives as inhibitors of HCV Replicon RNA replication. In particular, the invention is concerned with novel and known purine and pyrimidine nucleoside derivatives, their use as inhibitors of 5 subgenomic Hepatitis C Virus (HCV) RNA replication and pharmaceutical compositions of such compounds. For the novel purine and pyrimidine nucleoside derivatives the invention is also concerned with a process for their manufacture, pharmaceutical compositions and the use of such compounds in medicine. The compounds of this invention have potential use as therapeutic agents for the 10 treatment of HCV infections.

Hepatitis C virus is the leading cause of chronic liver disease throughout the world. Patients infected with HCV are at risk of developing cirrhosis of the liver and subsequent hepatocellular carcinoma and hence HCV is the major indication for liver transplantation. Only two approved therapies are currently available for 15 the treatment of HCV infection (R.G. Gish, Sem.Liver.Dis., 1999, 19, 35). These are interferon- α monotherapy and, more recently, combination therapy of the nucleoside analogue, ribavirin (Virazole), with interferon- α .

Ribavirin is a broad spectrum antiviral agent with activity against a range of DNA and RNA viruses (R.A.Smith and W. Kirkpatrick (Eds.): *Ribavirin – A Broad 20 Spectrum Antiviral Agent*, Academic Press, New York, 1980) but its mechanism of action has not been conclusively established and a number of distinct properties of ribavirin have been identified which may vary in relative importance for differing 25 viral disease conditions. These properties include mediation of the immune response (C. D. Hultgren et al, J.Gen.Viro., 1998, 79, 2381), lowering of serum alanine aminotransferase (ALT) levels (G. Dusheiko et al, J. Hepatol., 1996, 25, 591), inhibition as the monophosphate of inosine monophosphate dehydrogenase

- 2 -

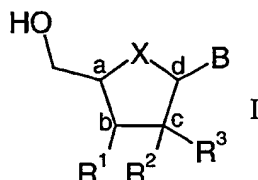
(IMPDH) (D.G.Streeter et al, Proc.Natl.Acad.Sci., 1973, 70,1174) and direct inhibition of viral DNA or RNA replication (R.W.Sidwell et al, Science, 177,705).

Many of the drugs approved for the treatment of viral infections are nucleosides or nucleoside analogues and most of these nucleoside analogue drugs inhibit viral replication, following conversion to the corresponding triphosphates, through inhibition of the viral polymerase enzymes. This conversion to the triphosphate is commonly mediated by cellular kinases and therefore the direct evaluation of nucleosides as inhibitors of HCV replication is only conveniently carried out using a cell-based assay. For HCV the availability of a true cell-based viral replication assay or animal model of infection is lacking.

Hepatitis C virus belongs to the family of Flaviridae. It is an RNA virus, the RNA genome encoding a large polyprotein which after processing produces the necessary replication machinery to ensure synthesis of progeny RNA. It is believed that most of the non-structural proteins encoded by the HCV RNA genome are involved in RNA replication. Lohmann et al. [V. Lohmann et al., Science, 1999, 285, 110-113] have described the construction of a Human Hepatoma (Huh7) cell line in which subgenomic HCV RNA molecules have been introduced and shown to replicate with high efficiency. It is believed that the mechanism of RNA replication in these cell lines is identical to the replication of the full length HCV RNA genome in infected hepatocytes. The subgenomic HCV cDNA clones used for the isolation of these cell lines have formed the basis for the development of a cell-based assay for identifying nucleoside analogue inhibitors of HCV replication.

The compounds of formula I have been shown to be inhibitors of subgenomic Hepatitis C Virus replication in a hepatoma cell line. These compounds have the potential to be efficacious as antiviral drugs for the treatment of HCV infections in human.

This object can be achieved by use of compounds of formula I



wherein

- 3 -

R^1 is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy, halogen, cyano, isocyano or azido;

R^2 is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine;

R^3 is hydrogen; or

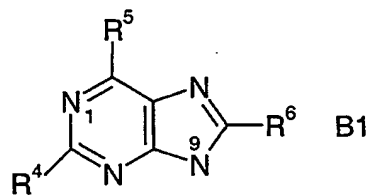
5 R^2 and R^3 together represent $=CH_2$; or

R^2 and R^3 represent fluorine;

X is O, S or CH_2 ;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

10 B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

R^4 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen or SH;

15 R^5 is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclamino, halogen, NR^7R^8 , $NHOR^9$, $NHNR^7R^8$ or SH;

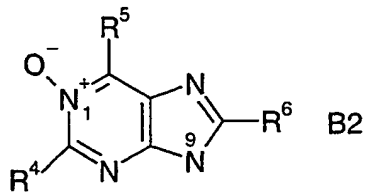
R^6 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen, SH or cyano;

20 R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R^9 is hydrogen, alkyl or aryl; or

B signifies an oxidised purine base B2 which is connected through the 9-nitrogen of formula

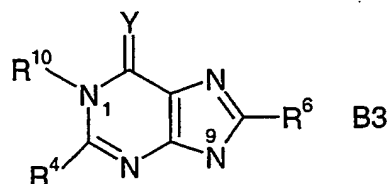
- 4 -



wherein

R⁴, R⁵ and R⁶ are as defined above; or

B signifies a purine base B3 which is connected through the 9-nitrogen of formula



5

wherein

R⁴ and R⁶ are as defined above;

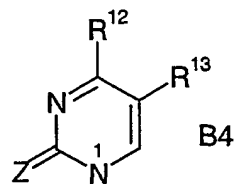
R¹⁰ is hydrogen, alkyl or aryl;

Y is O, S or NR¹¹;

10 R¹¹ is hydrogen, hydroxy, alkyl, OR⁹, heterocyclyl or NR⁷R⁸;

R⁷, R⁸ and R⁹ are as defined above; or

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



15 wherein

Z is O or S;

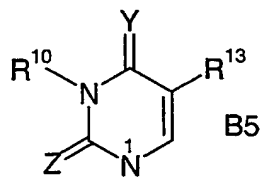
- 5 -

R^{12} is hydrogen, hydroxy, alkyl, alkoxy, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclamino, halogen, NR^7R^8 , $NHOR^9$, $NHNR^7R^8$ or SH;

R^{13} is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen;

R^7 , R^8 and R^9 are as defined above; or

- 5 B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula



wherein

Y , Z , R^{10} and R^{13} are as defined above

- 10 for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

The term "alkyl" as used herein denotes an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 12 carbon atoms, such as methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, pentyl, hexyl, heptyl, octyl, nonyl, decyl, undecyl, dodecyl including their isomers. Preferably, the term "alkyl" denotes an optionally substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms.

- Suitable substituents for the alkyl chain can be selected from one or more of
20 aryl, heterocyclyl, cycloalkyl,
nitro, cyano, azido,
amino, alkyl amino, dialkyl amino, cycloalkyl amino, aryl amino, diarylamino, heterocyclyl amino,
hydroxy, alkoxy, aryloxy, heterocyclyloxy, cycloalkoxy, thio, alkylthio, arylthio,
25 heterocyclthio,

alkyl carbonyl, cycloalkyl carbonyl, aryl carbonyl, heterocycl carbonyl, carboxy, alkoxy carbonyl, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, dialkylaminocarbonyl, diarylaminocarbonyl, heterocyclaminocarbonyl.

Aryl, heterocycl or cycloalkyl as substituents for the alkyl group can also be substituted with one or more methyl, ethyl, n-propyl, i-propyl, tert.-butyl, trifluoromethyl, hydroxy, methoxy, ethoxy, propyloxy, amino, alkylamino, arylamino, dialkylamino, diarylamino, heterocyclamino, vinyl, allyl, carboxy, alkylcarbonyl, alkoxy carbonyl, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, dialkylaminocarbonyl, diarylaminocarbonyl, heterocyclaminocarbonyl, fluorine, chlorine, bromine, iodine, cyano or nitro.

Alkyl in R¹ is preferably an unsubstituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms and most preferred methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl or pentyl.

Alkyl in R⁴ is preferably an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms. Suitable substituents for the alkyl group are selected from one or more of aryl or heterocycl as defined below. The aryl or heterocycl can also be alkylated with one or more methyl or ethyl or halogenated with fluorine, chlorine, bromine or iodine. Preferably alkyl in R⁴ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, phenylmethyl (benzyl), chlorphenylmethyl, phenylethyl, phenylpropyl, pyridylmethyl, chlorpyridylmethyl, pyridylethyl, pyridylpropyl, thienylmethyl, thienylethyl, thienylpropyl.

Alkyl in R⁵ is preferably an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms. Suitable substituents for the alkyl group are selected from one or more of aryl or heterocycl as defined below. The aryl or heterocycl can also be alkylated with one or more methyl or ethyl or halogenated with fluorine, chlorine, bromine or iodine. Preferably alkyl in R⁵ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, phenylmethyl (benzyl), chlorphenylmethyl, 1-phenylethyl, 2-phenylethyl, phenylpropyl, pyridylmethyl, chlorpyridylmethyl, pyridylethyl, pyridylpropyl, thienylmethyl, thienylethyl, thienylpropyl.

Alkyl in R⁶ is preferably an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms. Suitable substituents

for the alkyl group are selected from one or more of hydroxy, aryl or heterocyclyl as defined below. The aryl or heterocyclyl can also be alkylated with one or more methyl or ethyl or halogenated with fluorine, chlorine, bromine or iodine.

Preferably alkyl in R⁶ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, hydroxymethyl, 1-hydroxyethyl, 2-hydroxyethyl, hydroxypropyl, 1-hydroxy-1-methyl-ethyl, 2-hydroxy-2-methyl-ethyl, phenylmethyl (benzyl), chlorphenylmethyl, phenylethyl, phenylpropyl, pyridylmethyl, chlorpyridylmethyl, pyridylethyl, pyridylpropyl, thienylmethyl, thienylethyl, thienylpropyl.

Alkyl in R⁷ and R⁸ (for NR⁷R⁸) is independently of each other preferably an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 12 carbon atoms. Suitable substituents for the alkyl group are selected from one or more of aryl, heterocyclyl, cycloalkyl, nitro, amino, alkyl amino, dialkyl amino, cycloalkyl amino, aryl amino, heterocyclyl amino, alkyl carbonyl, cycloalkyl carbonyl, aryl carbonyl, heterocyclyl carbonyl. The aryl, heterocyclyl or cycloalkyl can also be substituted with one or more methyl, ethyl, n-propyl, i-propyl, tert.-butyl, trifluoromethyl, methoxy, ethoxy, propyloxy, amino, vinyl, allyl, carboxy, alkylcarbonyl, fluorine, chlorine, bromine, iodine or aminosulphonyl. Preferably alkyl in R⁷ and R⁸ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, pentyl, hexyl, heptyl, aminomethyl, aminoethyl, aminopropyl, aminobutyl, aminopentyl, chlormethyl, chlorethyl, chlorpropyl, cyanomethyl, cyanoethyl, cyanopropyl, phenylmethyl (benzyl), 1-phenylethyl, 2-phenylethyl, 1(S)-methyl-2-phenylethyl, 1(R)-methyl-2-phenylethyl, 1-phenylpropyl, 2-phenylpropyl, 3-phenylpropyl, phenylbutyl, phenylpentyl, phenylhexyl, 1-benzyl-1-methylethyl, chlorphenylmethyl, dichlorphenylmethyl, 2-chlorphenylethyl, 3-chlorphenylethyl, 4-chlorphenylethyl, dichlorphenylethyl, tolylmethyl, tolylethyl, tolylpropyl, tolylbutyl, methoxyphenylmethyl, methoxyphenylethyl, methoxyphenylpropyl, methoxyphenylbutyl, aminophenylmethyl, aminophenylethyl, aminophenylpropyl, aminophenylbutyl, phenolmethyl, phenoletethyl, phenolpropyl, phenolbutyl, naphthylmethyl, naphthylethyl, naphthylpropyl, naphthylbutyl, 2-pyridylmethyl, 3-pyridylmethyl, 4-pyridylmethyl, pyridylethyl, pyridylpropyl, methylpyridylmethyl, methylpyridylethyl, methylpyridylpropyl, chlorpyridylmethyl, chlorpyridylethyl, chlorpyridylpropyl, pyrrolmethyl, pyrrolethethyl, pyrrolpropyl, pyrrolbutyl, methylpyrrolmethyl, methylpyrrolethethyl, methylpyrrolpropyl, methylpyrrolylbutyl, imidazolylmethyl, imidazolethethyl, imidazolylpropyl, imidazolylbutyl, 2-(3-indolyl)methyl, 2-(3-indolyl)ethyl, 2-(3-indolyl)propyl,

5 morpholinylmethyl, morpholinylethyl, morpholinylpropyl, morpholinylbutyl,
thienylmethyl, thienylethyl, 2-(2-thienyl)ethyl, thietylpropyl, thietylbutyl,
cyclohexylmethyl, 1-cyclohexylethyl, 2-cyclohexylethyl, cyclohexylpropyl,
cyclohexylbutyl, 2-(4-cyanomethylphenyl)ethyl, 2-(3,4-dimethoxyphenyl)ethyl, 2-
(4-hydroxyphenyl)ethyl, (5-chloro-2-methoxyphenyl)methyl, (2-
methylphenyl)methyl, (3-methyl)butyl, 4-(aminophenyl)methyl, 2-(4-
morpholinyl)ethyl, 2(R,S)-phenylpropyl, 2-(4-Methylphenyl)ethyl, 2-(1-methyl-2-
pyrrolyl)ethyl, 2-(4-aminosulphonylphenyl)ethyl, 2-ethyl-4-imidazolyl, methyl-1-
naphthyl, 2-(4-chlorophenyl)ethyl, 2-(2,4-dichlorophenyl)ethyl, 4-fluorobenzyl, 4-
10 (hydroxycarbonyl)benzyl, 4-trifluoromethyl)benzyl, 2,5-dimethoxy)benzyl, 2-(2-
thienyl)ethyl, 2-(4-aminophenyl)ethyl, 2-Phenoxyethyl, (2-thienyl)methyl, 4-(tert-
Butyl)benzyl, 1(R)-Phenylethyl, 1(S)-Phenylethyl, 2-Hydroxy-1(S)-phenyl)ethyl.

15 Alkyl in R⁹ (for NHOR⁹) is preferably an unsubstituted or substituted
straight or branched chain hydrocarbon residue containing 1 to 12 carbon atoms
such as methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, pentyl, hexyl,
heptyl, octyl, nonyl, decyl, undecyl, dodecyl including their isomers. A suitable
substituent for the alkyl group is the aryl group as defined below. The aryl can also
be substituted with one or more methyl, ethyl, trifluoromethyl, methoxy, ethoxy,
hydroxy, amino, fluorine, chlorine, bromine or iodine. Preferred alkyl in R⁹ is
20 methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, pentyl, phenylmethyl
(benzyl), phenylethyl, phenylpropyl, phenylbutyl, chlorphenylmethyl,
chlorphenylethyl, tolylmethyl, tolylethyl, tolylpropyl, methoxyphenylmethyl,
methoxyphenylethyl, aminophenylmethyl, aminophenylethyl, phenolmethyl,
phenolethyl.

25 Alkyl in R¹⁰ is preferably an unsubstituted or substituted straight or branched
chain hydrocarbon residue containing 1 to 12 carbon atoms such as methyl, ethyl,
~~nonyl_isononyl_n-butyl_isobutyl_~~tert-butyl_n-pentyl_hexyl_hexyl_n-octyl_nonyl

Alkyl in R¹¹ is preferably an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms. A suitable substituent for the alkyl group is the aryl group as defined below. The aryl can also be substituted with one or more methyl, ethyl, trifluoromethyl, methoxy, ethoxy, 5 hydroxy, amino, fluorine, chlorine, bromine, iodine. Most preferred alkyl in R¹¹ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, pentyl, phenylmethyl (benzyl), phenylethyl, phenylpropyl, phenylbutyl, chlorphenylmethyl, chlorphenylethyl, tolylmethyl, tolylethyl, tolylpropyl, methoxyphenylmethyl, methoxyphenylethyl, aminophenylmethyl, aminophenylethyl, phenolmethyl, 10 phenoletethyl.

Alkyl in R¹² is preferably an unsubstituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms and most preferred methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl or pentyl.

Alkyl in R¹³ is preferably an unsubstituted or substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms such as methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl or pentyl, hexyl or heptyl. Suitable substituents for the alkyl group are selected from one or more of aryl, heterocycl, alkoxy or amino. The aryl or heterocycl can also be substituted with one or more methyl, trifluoromethyl, methoxy or amino. Preferably alkyl in R¹³ is methyl, ethyl, 15 propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, pentyl, hexyl, heptyl, methoxymethyl, ethoxymethyl, aminomethyl, aminoethyl, aminopropyl, aminobutyl, phenylmethyl (benzyl), phenylethyl, tolylmethyl, tolylethyl, 20 methoxyphenylmethyl, methoxyphenylethyl, aminophenylmethyl, aminophenylethyl, phenolmethyl, phenoletethyl, pyridylmethyl, pyridylethyl, methylpyridylmethyl, pyrrolylmethyl, pyrrolylethyl, methylpyrrolylmethyl, 25 methylpyrrolylethyl, imidazolylmethyl, imidazolylethyl, thienylmethyl, thienylethyl.

The term "cycloalkyl" as used herein denotes an optionally substituted cycloalkyl group containing 3 to 7 carbon atoms; e.g. cyclopropyl, cyclobutyl, 30 cyclopentyl, cyclohexyl or cycloheptyl, which can also be fused to an optionally substituted saturated, partially unsaturated or aromatic monocyclic, bicyclic or tricyclic heterocycle or carbocycle, e.g. to phenyl.

Suitable substituents for cycloalkyl can be selected from one or more of those named for alkyl.

- 10 -

Cycloalkyl in R⁵ is preferably an optionally substituted cycloalkyl group containing 3 to 7 carbon atoms, e.g. cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl or cycloheptyl. Suitable substituents for the cycloalkyl group are selected from aryl, heterocycl, cycloalkyl, hydroxy, nitro, halogen, amino, alkyl amino, dialkyl amino, cycloalkyl amino, aryl amino, heterocycl amino. The aryl or heterocycl can also be substituted with one or more of methyl, ethyl, trifluoromethyl, methoxy, amino, hydroxy, carboxy, fluorine, chlorine, bromine or iodine. Preferably cycloalkyl in R⁵ is cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl or cyclohexyl substituted with one or more aryl, heterocycl, methyl, amino, hydroxy, fluorine or chlorine.

Cycloalkyl in R⁷ and R⁸ (for NR⁷R⁸) is independently of each other preferably an optionally substituted cycloalkyl group containing 3 to 7 carbon atoms, e.g. cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl or cycloheptyl. Suitable substituents for the cycloalkyl group are selected from aryl, heterocycl, cycloalkyl, hydroxy, nitro, halogen, amino, alkyl amino, dialkyl amino, cycloalkyl amino, aryl amino, heterocycl amino. The aryl or heterocycl can also be substituted with one or more of methyl, ethyl, trifluoromethyl, methoxy, amino, hydroxy, carboxy, fluorine, chlorine, bromine or iodine. Preferably cycloalkyl in R⁷ and R⁸ is cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl or cyclohexyl substituted with one or more aryl, heterocycl, methyl, amino, hydroxy, fluorine or chlorine.

Cycloalkyl in R¹³ is preferably an optionally substituted cycloalkyl group containing 3 to 7 carbon atoms, e.g. cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl or cycloheptyl. Suitable substituents for the cycloalkyl group are selected from one or more of aryl, heterocycl, cycloalkyl, hydroxy, nitro, halogen, amino, alkyl amino, dialkyl amino, cycloalkyl amino, aryl amino or heterocycl amino. The aryl or heterocycl can also be substituted with one or more of methyl, ethyl, trifluoromethyl, methoxy, amino, hydroxy, carboxy, fluorine, chlorine, bromine or iodine. Preferably cycloalkyl in R¹³ is cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl, cycloheptyl or cyclohexyl substituted with one or more of aryl, heterocycl, methyl, amino, hydroxy, fluorine or chlorine.

The term "alkoxy" as used herein denotes an optionally substituted straight or branched chain alkyl-oxy group wherein the "alkyl" portion is as defined above such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, pentyloxy, hexyloxy, heptyloxy including their isomers.

Suitable substituents for the alkoxy group are selected from aryl, hydroxy, halogen or amino.

Alkoxy in R¹ is preferably an optionally substituted straight or branched chain alkyl-oxy group such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyoxy, i-butyloxy, tert.-butyloxy. Suitable substituents for the alkoxy group are selected from one ore more of aryl, halogen or amino. Preferably alkoxy in R¹ is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, phenylmethoxy, tolylmethoxy, fluormethoxy, chlormethoxy, bromomethoxy, fluorethoxy, chlorethoxy, bromomethoxy, aminomethoxy, aminoethoxy, aminopropyloxy.

Alkoxy in R² is preferably an optionally substituted straight or branched chain alkyl-oxy group such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyoxy, i-butyloxy, tert.-butyloxy. Suitable substituents for the alkoxy group are selected from one ore more of aryl, halogen or amino. Preferably alkoxy in R² is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, phenylmethoxy, tolylmethoxy, fluormethoxy, chlormethoxy, bromomethoxy, fluorethoxy, chlorethoxy, bromomethoxy, aminomethoxy, aminoethoxy, aminopropyloxy.

Alkoxy in R⁴ is preferably an optionally substituted straight or branched chain alkyl-oxy group such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyoxy, i-butyloxy, tert.-butyloxy. Suitable substituents for the alkoxy group are selected from one ore more of aryl, halogen or amino. Preferably alkoxy in R⁴ is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, phenylmethoxy, tolylmethoxy, fluormethoxy, chlormethoxy, bromomethoxy, fluorethoxy, chlorethoxy, bromomethoxy, aminomethoxy, aminoethoxy, aminopropyloxy.

Alkoxy in R⁵ is preferably an optionally substituted straight or branched chain alkyl-oxy group such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyoxy, i-butyloxy, tert.-butyloxy. Suitable substituents for the alkoxy group are selected from one ore more of aryl, halogen or amino. Preferably alkoxy in R⁵ is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, phenylmethoxy, tolylmethoxy, fluormethoxy, chlormethoxy, bromomethoxy, fluorethoxy, chlorethoxy, bromomethoxy, aminomethoxy, aminoethoxy, aminopropyloxy.

Alkoxy in R⁶ is preferably an optionally substituted straight or branched chain alkyl-oxy group such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy. Suitable substituents for the alkoxy group are selected from one or more of aryl, halogen or amino. Preferably alkoxy in R⁶ is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, phenylmethoxy, tolylmethoxy, fluoromethoxy, chlormethoxy, bromomethoxy, fluorethoxy, chlorethoxy, bromomethoxy, aminomethoxy, aminoethoxy, aminopropyloxy.

Alkoxy in R¹² is preferably an optionally substituted straight or branched chain alkyl-oxy group such as methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy. Suitable substituents for the alkoxy group are selected from one or more of aryl, halogen or amino. Preferably alkoxy in R¹² is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy, phenylmethoxy, tolylmethoxy, fluoromethoxy, chlormethoxy, bromomethoxy, fluorethoxy, chlorethoxy, bromomethoxy, aminomethoxy, aminoethoxy, aminopropyloxy.

The term "alkoxyalkyl" as used herein denotes an alkoxy group as defined above which is bonded to an alkyl group as defined above. Examples are methoxymethyl, methoxyethyl, methoxypropyl, ethoxymethyl, ethoxyethyl, ethoxypropyl, propyloxypropyl, methoxybutyl, ethoxybutyl, propyloxybutyl, butyloxybutyl, tert.-butyloxybutyl, methoxypentyl, ethoxypentyl, propyloxypentyl, butyloxypentyl, tert.-butyloxypentyl, pentyloxypentyl, methoxyhexyl, ethoxyhexyl, propyloxyhexyl, butyloxyhexyl, tert.-butyloxyhexyl, pentyloxyhexyl, hexyloxyhexyl, methoxyheptyl, ethoxyheptyl, propyloxyheptyl, butyloxyheptyl, tert.-butyloxyheptyl, pentyloxyheptyl, hexyloxyheptyl, heptyloxyheptyl including their isomers.

Alkoxyalkyl in R¹³ is preferably methoxymethyl, methoxyethyl, methoxypropyl, ethoxymethyl, ethoxyethyl, ethoxypropyl.

The term "alkenyl" as used herein denotes to unsubstituted or substituted hydrocarbon chain radical having from 2 to 7 carbon atoms, preferably from 2 to 4 carbon atoms, and having one or two olefinic double bonds, preferably one olefinic double bond. Examples are vinyl, 1-propenyl, 2-propenyl (allyl) or 2-but enyl (crotyl).

- 13 -

The term "alkenylalkyl" as used herein denotes an alkenyl group as defined above which is bonded to an alkyl group as defined above. Examples are vinylmethyl (e.g. 1-propenyl or 2-propenyl), 1-propenylmethyl, 2-propenylmethyl or 2-butenylmethyl.

5 Alkenylalkyl in R⁷ and R⁸ (for NR⁷R⁸) is independently of each other preferably 1-propenyl, 2-propenyl, 1-propenylmethyl or 2-propenylmethyl.

10 The term "alkynyl" as used herein denotes to unsubstituted or substituted hydrocarbon chain radical having from 2 to 7 carbon atoms, preferably 2 to 4 carbon atoms, and having one or where possible two triple bonds, preferably one triple bond. Examples are ethynyl, 1-propynyl, 2-propynyl, 1-butynyl, 2-butynyl or 15 3-butynyl.

The term "alkynylalkyl" as used herein denotes an alkynyl group as defined above which is bonded to an alkyl group as defined above. Examples are ethynylmethyl, 1-propynylmethyl, 2-propynylmethyl, 1-butynylmethyl, 2-butynylmethyl or 3-butynylmethyl.

15 Alkynylalkyl in R⁷ and R⁸ (for NR⁷R⁸) is independently of each other preferably ethynylmethyl, 1-propynylmethyl or 2-propynylmethyl.

20 The term "hydroxyalkyl" as used herein denotes a straight or branched chain alkyl group as defined above wherein 1, 2, 3 or more hydrogen atoms are substituted by a hydroxy group. Examples are hydroxymethyl, 1-hydroxyethyl, 2-hydroxyethyl, 1-hydroxypropyl, 2-hydroxypropyl, 3-hydroxypropyl, hydroxyisopropyl, hydroxybutyl, hydroxy-isobutyl, hydroxy-tert.-butyl, hydroxypentyl, hydroxyhexyl, hydroxyheptyl and the like.

25 Hydroxyalkyl in R¹, R⁷, R⁸, R¹³ is preferably hydroxymethyl, 1-hydroxyethyl, 2-hydroxyethyl, hydroxypropyl, hydroxy-isopropyl, hydroxybutyl, hydroxy-isobutyl, hydroxy-tert.-butyl, hydroxypentyl, hydroxyhexyl, hydroxyheptyl and preferred hydroxymethyl, 1-hydroxyethyl, 2-hydroxyethyl, 1-hydroxypropyl, 1-propanol, 2-propanol, 1-butanol, 2-butanol.

30 The term "haloalkyl" as used herein denotes a straight or branched chain alkyl group as defined above wherein 1, 2, 3 or more hydrogen atoms are substituted by a halogen. Examples are 1-fluoromethyl, 1-chloromethyl, 1-bromomethyl, 1-iodomethyl, trifluoromethyl, trichloromethyl, tribromomethyl,

triiodomethyl, 1-fluoroethyl, 1-chloroethyl, 1-bromoethyl, 1-idoethyl, 2-fluoroethyl, 2-chloroethyl, 2-bromoethyl, 2-idoethyl, 2,2-dichloroethyl, 3-bromopropyl or 2,2,2-trifluoroethyl and the like.

5 Haloalkyl in R⁵, R¹² and R¹³ is preferably 1-fluoromethyl, 1-chloromethyl, 1-bromomethyl, 1-iodomethyl, trifluoromethyl, trichloromethyl, tribromomethyl, triiodomethyl, 1-fluoroethyl, 1-chloroethyl, 1-bromoethyl, 1-idoethyl, 2-fluoroethyl, 2-chloroethyl, 2-bromoethyl, 2-idoethyl, 2,2-dichloroethyl, 3-bromopropyl or 2,2,2-trifluoroethyl.

10 The term "alkylthio" as used herein denotes a straight or branched chain (alkyl)S- group wherein the "alkyl" portion is as defined above and can be therefore as well substituted with substituents selected from one or more aryl or heterocycl. Examples are methylthio, ethylthio, n-propylthio, i-propylthio, n-butylthio, i-butylthio, tert.-butylthio, pentylthio, hexylthio, heptylthio, phenylmethylthio, phenylethylthio, phenylpropylthio, tolylmethylthio, tolylethylthio, tolylpropylthio, 15 pyridylmethylthio, pyridylethylthio, pyridpropylthio, pyrrolylmethylthio, pyrrolylethylthio or pyrrolylpropylthio.

20 Alkylthio in R⁴, R⁵, R⁶ and R¹² is preferably methylthio, ethylthio, n-propylthio, i-propylthio, n-butylthio, i-butylthio, tert.-butylthio, pentylthio, hexylthio, heptylthio, phenylmethylthio, phenylethylthio, phenylpropylthio, phenylbutylthio, tolylmethylthio, tolylethylthio, tolylpropylthio, pyridylmethylthio, pyridylethylthio, pyridpropylthio, pyrrolylmethylthio, pyrrolylethylthio or pyrrolylpropylthio. Preferred alkylthio in R⁴, R⁵, R⁶ and R¹² is methylthio, ethylthio, n-propylthio, i-propylthio, phenylmethylthio, phenylethylthio, phenylpropylthio, tolylmethylthio, tolylethylthio, pyridylmethylthio, 25 pyridylethylthio, pyrrolylmethylthio or pyrrolylethylthio.

30 The term "aryl" as used herein denotes an optionally substituted phenyl and naphthyl (e.g. 1-naphthyl, 2-naphthyl or 3-naphthyl), both optionally benz-fused to an optionally substituted saturated, partially unsaturated or aromatic monocyclic, bicyclic or tricyclic heterocycle or carbocycle e.g. to cyclohexyl or cyclopentyl such as 1,2-didehydronaphthyl, 1,2,3,4-tetradehydronaphthyl, anthryl, 1,2-didehydroanthryl, 1,2,3,4-tetradehydroanthryl, phenanthrenyl (e.g. 9-phenanthrenyl), 1,2-didehydrophenanthrenyl or 1,2,3,4-tetradehydrophenanthrenyl.

Suitable substituents for aryl can be selected from those named for alkyl, in addition however, halogen, hydroxy and optionally substituted alkyl, haloalkyl, alkenyl, alkynyl and aryloxy are substituents which can be added to the selection.

Examples for suitable aryls are tolyl, naphthyl (e.g. 1-naphthyl, 2-naphthyl or 3-naphthyl), p-ethylphenyl, p-propylphenyl, p-(i)propylphenyl, p-butylphenyl, p-(i)butylphenyl, p-(t)butylphenyl, 4-(2-methylpropyl)phenyl, p-hydroxyphenyl, p-fluorophenyl, p-chlorophenyl, p-bromophenyl, p-iodophenyl, p-methoxyphenyl, p-ethoxyphenyl, p-methylthiophenyl, p-perfluoromethylphenyl, p-perfluoromethoxyphenyl, biphenyl (e.g. 3-biphenylyl or 4-biphenylyl), p-phenoxyphenyl, m-ethylphenyl, m-propylphenyl, m-(i)propylphenyl, m-butylphenyl, m-(i)butylphenyl, m-(t)butylphenyl, m-hydroxyphenyl, m-fluorophenyl, m-chlorophenyl, m-bromophenyl, m-iodophenyl, m-methoxyphenyl, m-ethoxyphenyl, m-methylthiophenyl, m-perfluoromethylphenyl, m-perfluoromethoxyphenyl, m-phenoxyphenyl, o-ethylphenyl, o-propylphenyl, o-(i)propylphenyl, o-butylphenyl, o-(i)butylphenyl, o-(t)butylphenyl, o-hydroxyphenyl, o-fluorophenyl, o-chlorophenyl, o-bromophenyl, o-iodophenyl, o-methoxyphenyl, o-ethoxyphenyl, o-methylthiophenyl, p-methylthiophenyl, o-perfluoromethylphenyl, o-perfluoromethoxyphenyl or o-phenoxyphenyl. Aryl in R⁵ is preferably phenyl, naphthyl (e.g. 1-naphthyl, 2-naphthyl or 3-naphthyl), tolyl, phenanthrenyl (e.g. 9-phenanthrenyl), p-ethylphenyl, p-propylphenyl, p-(i)propylphenyl, p-butylphenyl, p-(i)butylphenyl, p-(t)butylphenyl, 4-(2-methylpropyl)phenyl, p-hydroxyphenyl, p-fluorophenyl, p-chlorophenyl, p-bromophenyl, p-iodophenyl, p-methoxyphenyl, p-ethoxyphenyl, p-methylthiophenyl, p-perfluoromethylphenyl, p-perfluoromethoxyphenyl, 3-biphenylyl, 4-biphenylyl, p-phenoxyphenyl, m-ethylphenyl, m-propylphenyl, m-(i)propylphenyl, m-butylphenyl, m-(i)butylphenyl, m-(t)butylphenyl, m-hydroxyphenyl, m-fluorophenyl, m-chlorophenyl, m-bromophenyl, m-iodophenyl, m-methoxyphenyl, m-ethoxyphenyl, m-methylthiophenyl, m-perfluoromethylphenyl, m-perfluoromethoxyphenyl, m-phenoxyphenyl, o-ethylphenyl, o-propylphenyl, o-(i)propylphenyl, o-butylphenyl, o-(i)butylphenyl, o-(t)butylphenyl, o-hydroxyphenyl, o-fluorophenyl, o-chlorophenyl, o-bromophenyl, o-iodophenyl, o-methoxyphenyl, o-ethoxyphenyl, o-methylthiophenyl, o-perfluoromethylphenyl, o-perfluoromethoxyphenyl or o-phenoxyphenyl.

Aryl in R⁵, R⁷, R⁸, R⁹, R¹⁰ and R¹² is preferably tolyl, p-ethylphenyl, p-hydroxyphenyl, p-fluorophenyl, p-chlorophenyl, p-bromophenyl, p-iodophenyl,

- 16 -

p-methoxyphenyl, p-ethoxyphenyl, p-perfluoromethylphenyl, p-perfluoromethoxyphenyl, 4-biphenyl, p-phenoxyphenyl, m-ethylphenyl, m-hydroxyphenyl, m-fluorophenyl, m-chlorophenyl, m-bromophenyl, m-iodophenyl, m-methoxyphenyl, m-perfluoromethylphenyl, m-perfluoromethoxyphenyl, m-phenoxyphenyl, o-ethylphenyl, o-hydroxyphenyl, o-fluorophenyl, o-chlorophenyl, o-bromophenyl, o-iodophenyl, o-methoxyphenyl, o-ethoxyphenyl, o-methylthiophenyl, o-perfluoromethylphenyl, o-perfluoromethoxyphenyl or o-phenoxyphenyl.

The term "aryloxy" as used herein denotes an aryl group as defined above which is bonded via an oxygen atom. Examples are phenoxy, naphthoxy and the like.

Aryloxy in R⁴, R⁵, R⁶ and R¹² is preferably phenoxy or naphthoxy, preferred phenoxy.

The term "arylthio" as used herein denotes an (aryl)S- group wherein the "aryl" portion is as defined above. Examples are phenylthio or naphthylthio.

Arylthio in R⁴, R⁵, R⁶ and R¹² is preferably phenylthio or naphthylthio, preferred phenylthio.

The term "heterocycl" as used herein denotes an optionally substituted saturated, partially unsaturated or aromatic monocyclic, bicyclic or tricyclic heterocyclic systems which contain one or more hetero atoms selected from nitrogen, oxygen and sulfur which can also be fused to an optionally substituted saturated, partially unsaturated or aromatic monocyclic carbocycle or heterocycle.

Examples of suitable heterocycles are oxazolyl, isoxazolyl, furyl, tetrahydrofuryl, 1,3-dioxolanyl, dihydropyranyl, 2-thienyl, 3-thienyl, pyrazinyl, isothiazolyl, isoquinolinyl, indolyl, didehydroindolyl, indazolyl, quinolinyl, dihydrooxazolyl, pyrimidinyl, benzofuranyl, tetrazolyl, 1-pyrrolidinyl, 2-pyrrolidinyl, 3-pyrrolidinyl, pyrrolidinonyl, (N-oxide)-pyridinyl, 1-pyrrolyl, 2-pyrrolyl, triazolyl e.g. 1,2,3-triazolyl or 1,2,4-triazolyl, 1-pyrazolyl, 2-pyrazolyl, 4-pyrazolyl, benzotriazolyl, piperidinyl, morpholinyl (e.g. 4-morpholinyl), thiomorpholinyl (e.g. 4-thiomorpholinyl), thiazolyl, pyridinyl, dihydrothiazolyl, imidazolidinyl, pyrazolinyl, benzothienyl, piperazinyl, 1-imidazolyl, 2-imidazolyl, 4-imidazolyl, thiadiazolyl e.g. 1,2,3-thiadiazolyl, 1,2,3,4-tetrahydroquinoline, 1,2,3,4-tetrahydroisoquinoline, benzothiazolyl, thianthrene (e.g. 1-thianthrenyl) or

heptamethyleneimine, 1,2,4,5-tetrahydro-3H-benzazepin-3-yl, 1,2,3,4-tetrahydro-2-isoquinolyl, 4-methylpiperazinyl, 1,3,4,5-tetrahydro-2H-benzazepin-2-yl, 2,3-dihydro-1-indolyl, 2-isoindolinyl, 2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl, 2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl, 8-amino sulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl, 7-amino sulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl, 10,11-dihydro-5H-dibenzo[a,d]cyclohepten-5-yl, 1-hexamethyleneimino, 4-hydroxypiperidin-1-yl, 1,2,3,4-tetrahydro-2-isoquinolyl, 4-phenyl-1-piperazinyl.

Suitable substituents for heterocyclyl can be selected from those named for alkyl, in addition however, optionally substituted alkyl, alkenyl, alkynyl, an oxo group (=O) or aminosulphonyl are substituents which can be added to the selection.

Heterocyclyl in R⁴ is preferably unsubstituted or substituted furyl, tetrahydrofuryl, thienyl, indolyl, indazolyl, pyrimidinyl, benzofuranyl, 1-pyrrolidinyl, pyrrolidinonyl, (N-oxide)-pyridinyl, pyrrolyl, piperidinyl, morpholinyl, imidazolyl or benzothiazolyl. Suitable substituents for heterocyclyl in R⁴ can be selected from unsubstituted or substituted alkyl, unsubstituted or substituted aryl, nitro, cyano and amino.

Heterocyclyl in R⁵ is preferably unsubstituted or substituted oxazolyl, isoxazolyl, furyl, tetrahydrofuryl, 1,3-dioxolanyl, dihydropyranyl, thienyl, pyrazinyl, isothiazolyl, isoquinolinyl, 1-indolyl, didehydroindolyl, indazolyl, quinolinyl, dihydrooxazolyl, pyrimidinyl, benzofuranyl, tetrazolyl, 1-pyrrolidinyl, pyrrolidinonyl, (N-oxide)-pyridinyl, 1,2,3,6-tetradehydropyridine, 1-pyrrolyl, 2-pyrrolyl, triazolyl e.g. 1,2,4-triazolyl, 1-pyrazolyl, 2-pyrazolyl, benzotriazolyl, piperidinyl, 4-morpholinyl, 4-thiomorpholinyl, thiazolyl, pyridinyl, dihydrothiazolyl, imidazolidinyl, pyrazolinyl, benzothienyl, piperazinyl, 1-imidazolyl, thiadiazolyl e.g. 1,2,3-thiadiazolyl, benzothiazolyl, 1-thianthrenyl or heptamethyleneimine, 1,2,4,5-tetrahydro-3H-benzazepin-3-yl, 1,2,3,4-tetrahydro-2-isoquinolyl, 4-methylpiperazinyl, 1,3,4,5-tetrahydro-2H-benzazepin-2-yl, 2,3-dihydro-1-indolyl, 2-isoindolinyl, 2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl, 2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl, 8-amino sulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl, 7-amino sulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl, 10,11-dihydro-5H-dibenzo[a,d]cyclohepten-5-yl, 1-hexamethyleneimino, 4-hydroxypiperidin-1-yl, 1,2,3,4-tetrahydro-2-isoquinolyl, 4-phenyl-1-piperazinyl.

Suitable substituents for heterocycl in R⁵ can be selected from unsubstituted or substituted alkyl as defined above, unsubstituted or substituted aryl as defined above, nitro, cyano and amino. Examples for substituted heterocycl are methylpiperazinyl, ethylpiperazinyl, propylpiperazinyl, butylpiperazinyl, phenylpiperazinyl, methoxyphenylpiperazinyl (e.g. 4-(2-Methoxyphenyl)piperazinyl), ethoxyphenylpiperazinyl, propyloxyphenylpiperazinyl, benzo-fused thianthrene or 4-(4-Fluorophenyl)-1,2,5,6-tetrahydropyridyl.

Heterocycl in R⁶ is preferably unsubstituted or substituted oxazolyl, 10 isoxazolyl, furyl, tetrahydrofuryl, 1,3-dioxolanyl, dihydropyranyl, 2-thienyl, 3-thienyl, pyrazinyl, isothiazolyl, isoquinolinyl, indolyl, didehydroindolyl, indazolyl, quinolinyl, dihydrooxazolyl, pyrimidinyl, benzofuranyl, tetrazolyl, 1-pyrrolidinyl, 2-pyrrolidinyl, 3-pyrrolidinyl, pyrrolidinonyl, (N-oxide)-pyridinyl, 1,2,3,6-tetradehydropyridine, pyrrolyl, 1,2,3-triazolyl, 1,2,4-triazolyl, 1-pyrazolyl, 2-pyrazolyl, 4-pyrazolyl, benzotriazolyl, 1-piperidinyl, 4-morpholinyl, thiomorpholinyl, thiazolyl, pyridinyl, dihydrothiazolyl, imidazolidinyl, 1-imidazolyl, 2-imidazolyl, 4-imidazolyl, pyrazolinyl, benzothienyl, piperazinyl, imidazolyl, thiadiazolyl e.g. 1,2,3-thiadiazolyl, 1,2,3,4-tetrahydroquinoline, 1,2,3,4-tetrahydroisoquinoline, benzothiazolyl, thianthrene or heptamethyleneimine.

20 Suitable substituents for heterocycl in R⁶ can be selected from unsubstituted or substituted alkyl as defined above, unsubstituted or substituted aryl as defined above, nitro, cyano and amino. Examples for substituted heterocycl are methylpiperazinyl, ethylpiperazinyl, propylpiperazinyl, butylpiperazinyl, phenylpiperazinyl, methoxyphenylpiperazinyl, ethoxyphenylpiperazinyl, propyloxyphenylpiperazinyl or benzo-fused thianthrene.

Heterocycl in R¹¹ or R¹² is preferably unsubstituted or substituted furyl, 30 tetrahydrofuryl, thienyl indolyl, indazolyl, pyrimidinyl, benzofuranyl, pyrrolidinyl, pyrrolidinonyl, (N-oxide)-pyridinyl, 1-pyrrolyl, piperidinyl, morpholinyl, imidazolyl or benzothiazolyl. Suitable substituents for heterocycl in R⁴ can be selected from unsubstituted or substituted alkyl, unsubstituted or substituted aryl, nitro, cyano and amino.

The term "heterocyclamino" refers to a group of formula (heterocycl)N(H), wherein heterocycl is as defined above. Examples are

furylamino, tetrahydrofurylamino, dihydropyranylamino, thienylamino,
pyrazinylamino, indolylamino, indazolylamino, quinolinylamino,
benzofuranylamin, pyrrolidinylamino, pyrrolidinonylamino, (N-oxide)-
pyridinylamino, pyrrolylamino, pyrazolylamino, benzotriazolylamino,
5 piperidinylamino, morpholinylamino, thiazolylamino, pyridinylamino,
imidazolidinylamino, benzothienylamino, imidazolylamino or
benzothiazolylamino.

Heterocyclylamino in R⁵ or R¹² is preferably furylamino,
tetrahydrofurylamino, dihydropyranylamino, thienylamino, pyrazinylamino,
10 indolylamino, indazolylamino, quinolinylamino, benzofuranylamin,
pyrrolidinylamino, pyrrolidinonylamino, (N-oxide)-pyridinylamino,
pyrrolylamino, pyrazolylamino, benzotriazolylamino, piperidinylamino,
morpholinylamino, thiazolylamino, pyridinylamino, imidazolidinylamino,
benzothienylamino, imidazolylamino or benzothiazolylamino.

15 The term "acyl" as used herein denotes a group of formula C(=O)R wherein
R is hydrogen, an unsubstituted or substituted straight or branched chain
hydrocarbon residue containing 1 to 7 carbon atoms or a phenyl group. Most
preferred acyl groups are those wherein R is hydrogen, an unsubstituted straight
chain or branched hydrocarbon residue containing 1 to 4 carbon atoms or a
20 phenyl group.

Acyl in R⁷ and R⁸ (for NR⁷R⁸) is independently of each other preferably
methylcarbonyl (acetyl), ethylcarbonyl (propionyl), propylcarbonyl, butylcarbonyl
or phenylcarbonyl (benzoyl).

25 The term halogen stands for fluorine, chlorine, bromine or iodine, preferable
fluorine, chlorine, bromine.

Halogen in R¹ is preferably fluorine, chlorine or iodine and more preferred
fluorine.

Halogen in R⁴ is preferably chlorine.

Halogen in R⁵ is preferably chlorine.

30 Halogen in R⁶ is preferably chlorine or bromine.

- 20 -

Halogen in R¹² or R¹³ is preferably fluorine, chlorine , bromine or iodine, more preferred fluorine, chlorine or bromine

Within the invention the term "X" represents O, S or CH₂, preferably O or CH₂. Most preferred "X" represents O.

5 Within the invention the term "Y" represents O, S or NR¹¹, wherein R¹¹ represents hydrogen, hydroxy or alkyl which denotes an unsubstituted or aryl-substituted straight or branched chain hydrocarbon residue containing 1 to 7 carbon atoms. Preferably "Y" represents O, S or NR¹¹ wherein R¹¹ represents hydrogen, hydroxy, phenylmethyl (benzyl), phenylethyl, phenylpropyl,
10 phenylbutyl.

Within the invention the term "Z" represents O or S, more preferred O.

In the pictorial representation of the compounds given throughout this application, a thickened tapered line (▼) indicates a substituent which is above the plane of the ring to which the asymmetric carbon belongs, a dotted line (---) indicates a substituent which is below the plane of the ring to which the
15 asymmetric carbon belongs, and a wavy line (~) indicates a substituent which can be either above or below the plane of the molecule. It is to be understood that the pictorial representation of the compounds given throughout the specification are set forth for convenience and are to be construed as inclusive of other forms including stereoisomers, enantiomers and racemates and are not to be construed as limited to the particular form shown.
20

Compounds of formula I exhibit stereoisomerism. The compounds of this invention can be any isomer of the compound of formula I or mixtures of these isomers. The compounds and intermediates of the present invention having one or
25 more asymmetric carbon atoms may be obtained as racemic mixtures of stereoisomers which can be resolved, at the appropriate steps in the process of this invention by methods known in the art to obtain a given stereoisomer or pure enantiomer having a desired stereoconfiguration. Alternatively, the desired isomers may be directly synthesised by methods known in the art.

30 Asymmetric carbon atoms in the compounds of the present invention are denoted as a, b, c and d. The stereoconfiguration of each of the asymmetric carbon atoms denoted as a, b, c, and d can be designated according to the particular stereoisomer it represents. Compounds of the present invention include those

compounds wherein the carbon atom denoted as "a" has the S, R, or R,S-configuration; the carbon atom denoted as "b" has the S, R, or R,S-configuration; the carbon atom denoted as "c" has the S, R, or R,S-configuration; and the carbon atom denoted as "d" has the S, R, or R,S-configuration. In a preferred embodiment 5 of the invention a, b, c and d denoting asymmetric carbon atoms and forming a α -D, β -D, α -L or β -L ribofuranosyl ring. Preferably a, b, c and d denoting asymmetric carbon atoms and forming an α -D or β -D ribofuranosyl ring and most preferred, β -D ribofuranosyl ring.

Compounds of formula I exhibit tautomerism that means that the 10 compounds of this invention can exist as two or more chemical compounds that are capable of facile interconversion. In many cases it merely means the exchange of a hydrogen atom between two other atoms, to either of which it forms a covalent bond. Tautomeric compounds exist in a mobile equilibrium with each other, so that attempts to prepare the separate substances usually result in the 15 formation of a mixture that shows all the chemical and physical properties to be expected on the basis of the structures of the components.

The most common type of tautomerism is that involving carbonyl, or keto, 20 compounds and unsaturated hydroxyl compounds, or enols. The structural change is the shift of a hydrogen atom between atoms of carbon and oxygen, with the rearrangement of bonds as indicated.

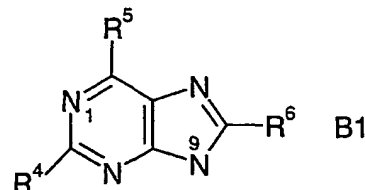
For example, in many aliphatic aldehydes and ketones, such as acetaldehyde, the keto form is the predominant one; in phenols, the enol form is the major component. An intermediate situation is represented for example in ethyl acetacetate, which at room temperature contains about 92.4 percent keto and 7.6 25 percent enol; at -78° C, the interconversion of the two forms is slow enough for the individual substances to be isolated.

It will be appreciated that within the present invention compounds of formula I exist in various tautomeric forms and that they are encompassed by the present invention.

- 22 -

A preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula

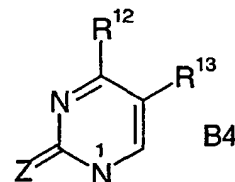


5 wherein

R⁴, R⁵, R⁶, R⁷, R⁸ and R⁹ are as defined in formula I;

with the proviso that R⁴ is not NH₂ and R⁵ is not NH(CH₃); or

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



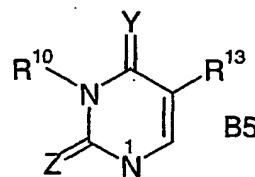
10

wherein

Z, R⁷, R⁸, R⁹, R¹², R¹³ are as defined in formula I;

with the proviso that R¹² is not hydroxy, alkoxy, N(CH₃)₂, N(H)NH(CH₃) or N(H)NH₂ and R¹³ is not hydroxyalkyl, chlorine or bromine; or

15 B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula



wherein

- 23 -

Y, Z, R¹⁰ and R¹³ are as defined in formula I;

with the proviso that R¹⁰ is not methyl or hydroxyethyl;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

5

A further preferred embodiment of the invention is the use of compounds of formula I wherein

R¹ is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy or halogen,

preferably wherein

10 R¹ is hydroxy;

R² is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine,

preferably wherein

R² is hydroxy;

R³ is hydrogen; or

15 R² and R³ represent fluorine;

X is O;

a, b, c and d denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring,

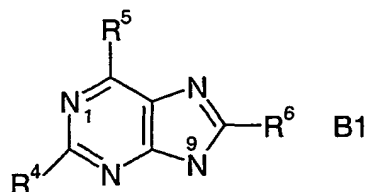
preferably wherein

20 a, b, c and d denoting asymmetric carbon atoms and forming a β -D-ribofuranosyl ring;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

A particularly preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



5 wherein

R⁴ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen or SH,

preferably wherein

R⁴ is hydrogen, chlorine or NH₂,

10 most preferred wherein

R⁴ is hydrogen;

R⁵ is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH,

15 preferably wherein

R⁵ is hydroxy, alkylthio, aryl, heterocycl, halogen, NR⁷R⁸ or SH,

most preferred wherein

R⁵ is alkylthio, aryl, heterocycl, halogen or NR⁷R⁸;

20 R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen, SH or cyano,

preferably wherein

R⁶ is hydrogen, halogen, heterocycl or NR⁷R⁸,

most preferred wherein

R^6 is hydrogen or halogen;

R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl,

5 preferably wherein

R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, alkenylalkyl or alkynylalkyl,

most preferred wherein

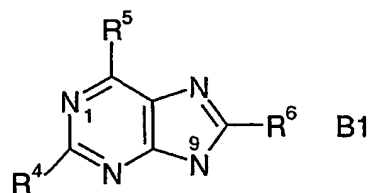
10 R^7 and R^8 are independently of each other hydrogen, alkyl, alkenylalkyl or alkynylalkyl;

R^9 is hydrogen, alkyl or aryl;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

15 A further preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

20 R^4 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen or SH,

preferably wherein

R⁴ is hydrogen or chlorine,

most preferred wherein

R⁴ is hydrogen;

5 R⁵ is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH,

preferably wherein

R⁵ is hydroxy, alkylthio, aryl, heterocycl, halogen, NR⁷R⁸ or SH,

most preferred wherein

10 R⁵ is alkylthio, aryl, heterocycl, halogen or NR⁷R⁸;

R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen, SH or cyano,

preferably wherein

R⁶ is hydrogen, halogen, heterocycl or NR⁷R⁸,

15 most preferred wherein

R⁶ is hydrogen or halogen;

R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl,

preferably wherein

20 R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, alkenylalkyl or alkynylalkyl;

R⁹ is hydrogen, alkyl or aryl;

with the proviso that R⁴ is not NH₂ and R⁵ is not NH(CH₃),

preferably

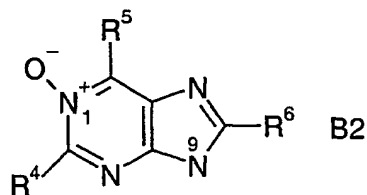
- 27 -

with the proviso that R⁵ is not NH(CH₃);

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

5 A particularly preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies an oxidised purine base B2 which is connected through the 9-nitrogen of formula



10 wherein

R⁴ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR⁷R⁸, halogen or SH,

preferably wherein

R⁴ is hydrogen;

15 R⁵ is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclylamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH,

preferably wherein

R⁵ is hydrogen, alkyl, heterocyclyl or NR⁷R⁸;

20 R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR⁷R⁸, halogen, SH or cyano,

preferably wherein

R⁶ is hydrogen;

- 28 -

R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl,

preferably wherein

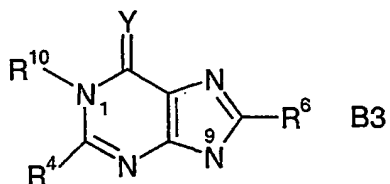
5 R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R⁹ is hydrogen, alkyl or aryl;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

10 Another preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a purine base B3 which is connected through the 9-nitrogen of formula



wherein

15 R⁴ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen or SH,

preferably wherein

R⁴ is hydrogen, NR⁷R⁸ or hydroxy;

20 R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen, SH or cyano,

preferably wherein

R⁶ is hydrogen, halogen or NR⁷R⁸;

- 29 -

R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl,

preferably wherein

R^7 and R^8 are independently of each other hydrogen or alkyl;

5 R^9 is hydrogen, alkyl or aryl;

R^{10} is hydrogen, alkyl or aryl,

preferably wherein

R^{10} is hydrogen or alkyl;

Y is O, S or NR^{11} ,

10 preferably wherein

Y is O, S, NH or N-alkyl;

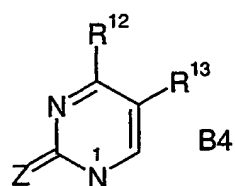
R^{11} is hydrogen, hydroxy, alkyl, OR^9 , heterocyclyl or NR^7R^8 ;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

15

Another preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



20

wherein

Z is O or S,

preferably wherein

Z is O;

R¹² is hydrogen, hydroxy, alkyl, alkoxy, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocycll, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH,

5 preferably wherein

R¹² is hydroxy, alkyl, heterocycll, NR⁷R⁸, NHOR⁹, heterocyclamino, NHNR⁷R⁸ or SH,

most preferred wherein

R¹² is hydroxy, alkyl or NR⁷R⁸;

10 R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen,

preferably wherein

R¹³ is hydrogen, alkyl or halogen,

most preferred wherein

R¹³ is hydrogen;

15 R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl,

preferably wherein

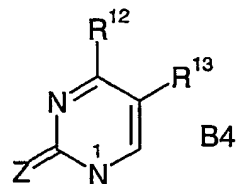
R⁷ and R⁸ are independently of each other hydrogen or alkyl;

R⁹ is hydrogen, alkyl or aryl;

20 for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

A further preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

5 Z is O or S,

preferably wherein

Z is O;

R¹² is hydrogen, alkyl, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclylamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH,

10 preferably wherein

R¹² is alkyl, heterocyclyl, NR⁷R⁸, NHOR⁹, heterocyclylamino, NHNR⁷R⁸ or SH,

most preferred wherein

R¹² is hydroxy, alkyl or NR⁷R⁸;

R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen,

15 preferably wherein

R¹³ is hydrogen, alkyl or halogen,

most preferred wherein

R¹³ is hydrogen;

20 R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl,

preferably wherein

- 32 -

R^7 and R^8 are independently of each other hydrogen or alkyl;

R^9 is hydrogen, alkyl or aryl;

with the proviso that R^{12} is not $N(CH_3)_2$, $N(H)NH(CH_3)$ or $N(H)NH_2$ and R^{13} is not hydroxyalkyl, chlorine or bromine,

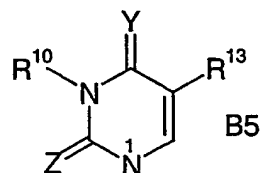
5 preferably

with the proviso that R^{12} is not $N(CH_3)_2$, $N(H)NH(CH_3)$ or $N(H)NH_2$;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

10 Another preferred embodiment of the invention is the use of compounds of formula I wherein

B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula



15 wherein

Y is O, S or NR^{11} ,

preferably wherein

Y is O or NR^{11} ;

Z is O or S,

20 preferably wherein

Z is O;

R^{10} is hydrogen, alkyl or aryl,

preferably wherein

R^{10} is hydrogen;

R^{13} is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen,
preferably wherein

R^{13} is hydrogen, alkyl or halogen;

- 5 for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

A further preferred embodiment of the invention is the use of compounds of formula I wherein

- 10 R^1 is hydrogen, halogen, hydroxy, alkyl, alkoxy, cyano or azido,

preferably wherein

R^1 is hydrogen, fluorine, hydroxy, C_{1-4} -alkyl, C_{1-4} -alkoxy, cyano or azido;

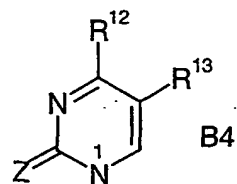
R^2 is hydrogen or hydroxy; or

R^2 and R^3 represent fluorine;

- 15 X is O or CH_2 ;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



20

wherein

Z is O;

- 34 -

R¹² is NR⁷R⁸;

R¹³ is hydrogen, alkyl or halogen,

preferably wherein

R¹³ is hydrogen, C₁₋₄-alkyl or fluorine;

5 R⁷ and R⁸ are independently of each other hydrogen or alkyl,

preferably wherein

R⁷ and R⁸ are independently of each other hydrogen or C₁₋₄-alkyl;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

10

Another preferred embodiment of the invention is the use of compounds of formula I wherein

R¹ is hydrogen, halogen, hydroxy, alkyl, alkoxy, cyano or azido;

preferably wherein

15 R¹ is hydrogen, fluorine, hydroxy, C₁₋₄-alkyl, C₁₋₄-alkoxy, cyano or azido;

R² is hydrogen or hydroxy; or

R² and R³ represent fluorine;

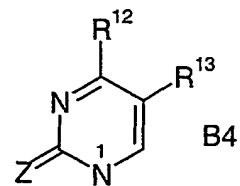
X is O or CH₂,

preferably wherein

20 X is CH₂;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

5 Z is O;

R¹² is NR⁷R⁸;

R¹³ is hydrogen, alkyl or halogen,

preferably wherein

R¹³ is hydrogen, C₁₋₄-alkyl or fluorine;

10 R⁷ and R⁸ are independently of each other hydrogen or alkyl,

preferably wherein

R⁷ and R⁸ are independently of each other hydrogen or C₁₋₄-alkyl;

with the proviso that R¹² is not N(CH₃)₂ and R¹³ is not chlorine or bromine,

preferably

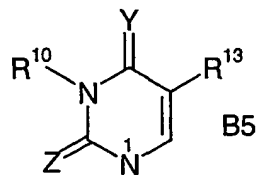
15 with the proviso that R¹² is not N(CH₃)₂;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

20 A further preferred embodiment of the invention is the use of compounds of formula I wherein

- 36 -

B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula



wherein

5 Y is O, S or NR¹¹;

Z is O or S;

R¹⁰ is hydrogen, alkyl or aryl;

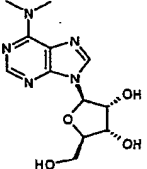
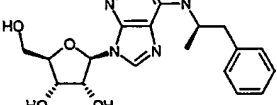
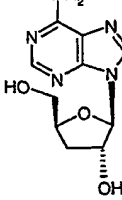
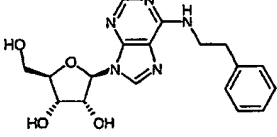
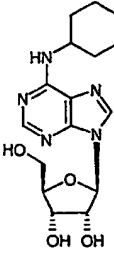
R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen;

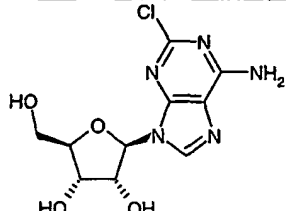
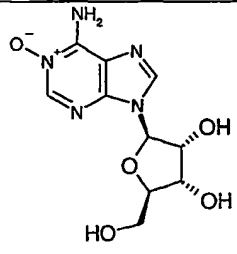
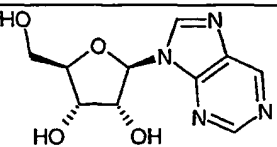
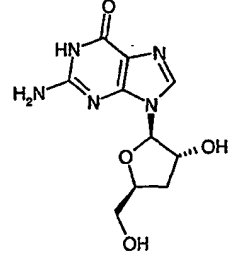
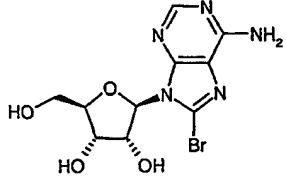
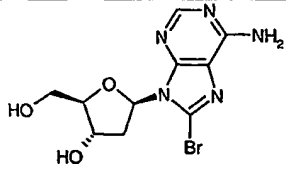
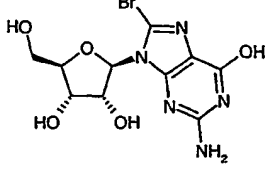
with the proviso that R¹⁰ is not methyl or hydroxyethyl;

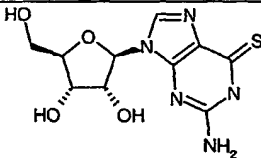
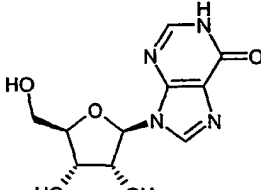
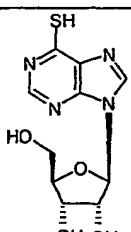
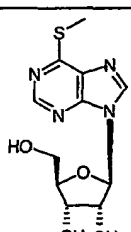
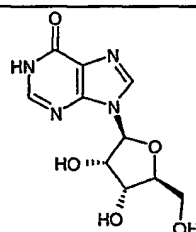
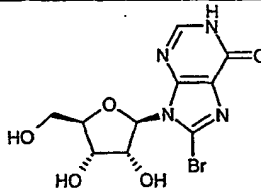
10 for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

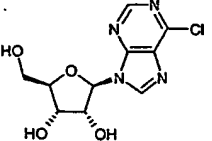
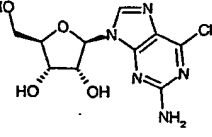
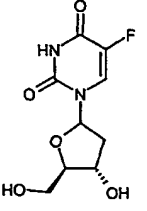
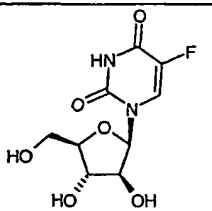
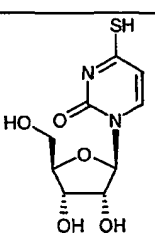
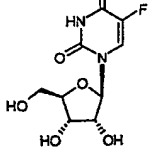
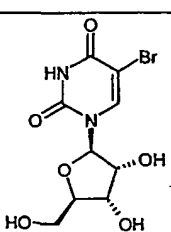
More preferred embodiments for the use of compound of formula I for the treatment of diseases mediated by the Hepatitis C Virus or for the preparation of a medicament for such treatment are set out in table 1 (see below):

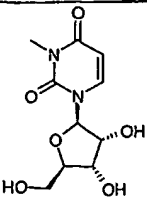
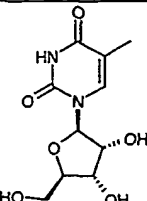
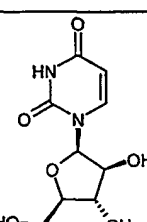
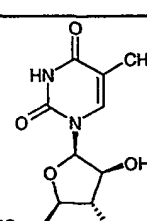
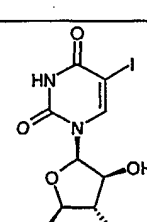
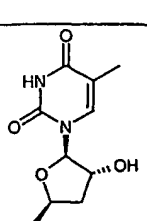
Table 1

Example	STRUCTURE	Name
1		6-Dimethylamino-9-(β-D-ribofuranosyl)purine
2		6-[1(S)-Methyl-2-phenylethylamino]-9-(β-D-ribofuranosyl)purine
3		3'-Deoxyadenosine
4		6-(Phenylethylamino)-9-(β-D-ribofuranosyl)purine
5		6-(Cyclohexylamino)-9-(β-D-ribofuranosyl)purine

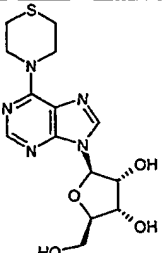
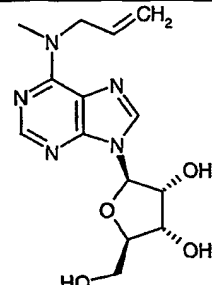
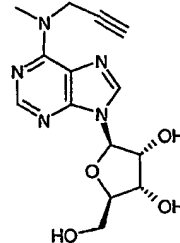
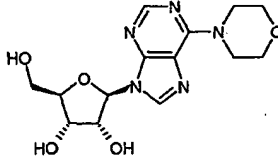
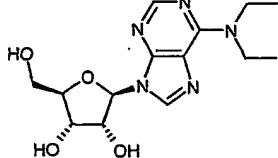
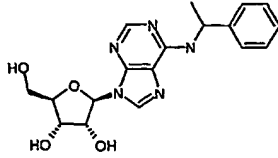
6		2-Chloroadenosine
7		Adenosine-1-oxide
8		9-(β-D-Ribofuranosyl)purine
9		3'-Deoxyguanosine
10		8-Bromoadenosine
11		8-Bromo-2'-deoxyadenosine
12		8-Bromoguanosine

13		6-Thioguanosine
14		Inosine
15		6-Thioinosine
16		6-Methylthio-9-(β -D-ribofuranosyl)purine
17		L-Inosine
18		8-Bromoinosine

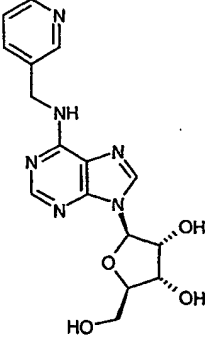
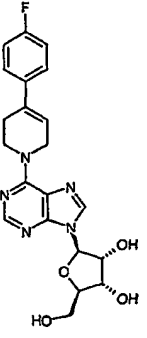
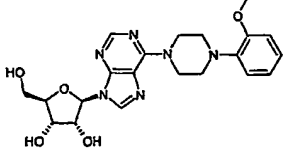
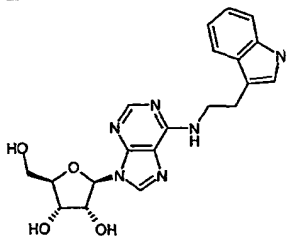
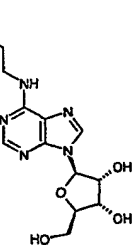
19		6-Chloro-9-(β -D-ribofuranosyl)purine
20		2-Amino-6-chloro-9-(β -D-ribofuranosyl)purine
21		2'-Deoxy-5-fluorouridine
22		1-(β -D-Arabinofuranosyl)-5-fluorouracil
23		4-Thiouridine
24		5-Fluorouridine
25		5-Bromouridine

26		3-Methyluridine
27		5-Methyluridine
28		1-(β -D-Arabinofuranosyl)uracil
29		1-(β -D-Arabinofuranosyl)-5-methyluracil
30		1-(β -D-Arabinofuranosyl)-5-iodouracil
31		3'-Deoxy-5-methyluridine

32		5-Fluorocytidine
33		1-(β-D-Arabinofuranosyl)-5-fluorocytosine
34		5-Methylcytidine
35		2',3'-Dideoxycytidine
36		N4-Acetylcytidine
37		3'-Deoxycytidine
38		6-(N-Methylpropylamino)-9-(β-D-ribofuranosyl)purine

39		9-(β -D-Ribofuranosyl)-6-(4-thiomorpholinyl)purine
40		6-(N-Methyl-2-propenylamino)-9-(β -D-ribofuranosyl)purine
41		6-(N-Methyl-2-propynylamino)-9-(β -D-ribofuranosyl)purine
42		6-(4-Morpholinyl)-9-(β -D-ribofuranosyl)purine
43		6-Diethylamino-9-(β -D-ribofuranosyl)purine
44		6-(1(R,S)-Phenylethylamino)-9-(β -D-ribofuranosyl)purine

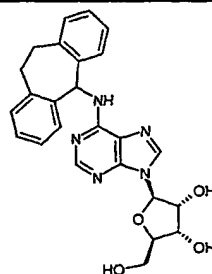
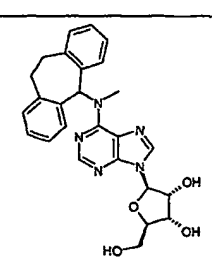
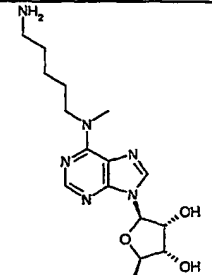
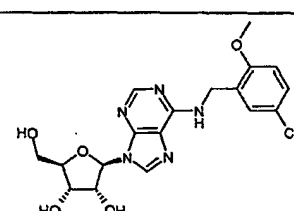
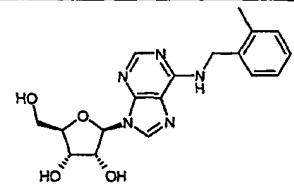
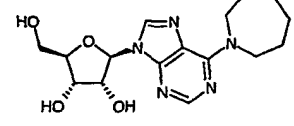
45		6-(1-Benzyl-1-methylethylamino)-9-(β -D-ribofuranosyl)purine
46		6-(3-Phenylpropylamino)-9-(β -D-ribofuranosyl)purine
47		9-(β -D-Ribofuranosyl)-6-[2-(2-thienyl)ethylamino]purine
48		6-Dibenzylamino-9-(β -D-ribofuranosyl)purine
49		6-Hexylamino-9-(β -D-ribofuranosyl)purine

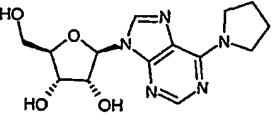
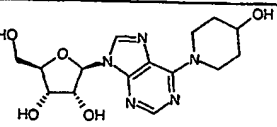
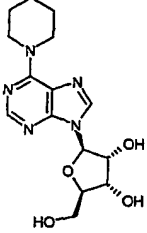
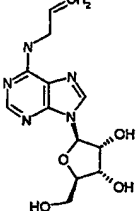
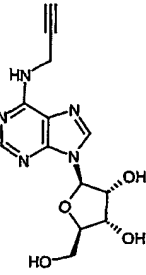
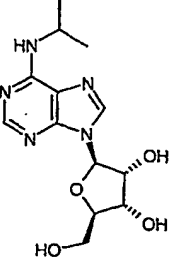
50		6-(3-Pyridylmethylamino)-9-(β -D-ribofuranosyl)purine
51		6-[4-(4-Fluorophenyl)-1,2,5,6-tetrahydropyridyl]-9-(β -D-ribofuranosyl)purine
52		6-[4-(2-Methoxyphenyl)piperazinyl]-9-(β -D-ribofuranosyl)purine
53		6-[2-(3-Indolyl)ethylamino]-9-(β -D-ribofuranosyl)purine
54		6-[2-(4-Chlorophenyl)ethylamino]-9-(β -D-ribofuranosyl)purine

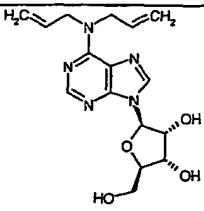
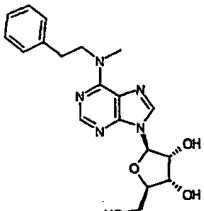
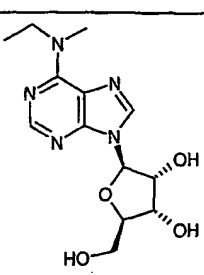
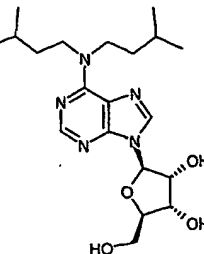
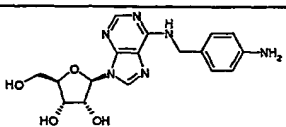
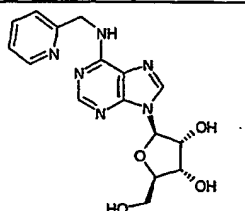
55		6-(N-Methylphenylamino)-9-(β -D-ribofuranosyl)purine
56		9-(β -D-Ribofuranosyl)-6-(1,2,4,5-tetrahydro-3H-benzazepin-3-yl)purine
57		9-(β -D-Ribofuranosyl)-6-(1,2,3,4-tetrahydro-2-isoquinolyl)purine
58		6-(4-Methylpiperazinyl)-9-(β -D-ribofuranosyl)purine
59		9-(β -D-Ribofuranosyl)-6-(1,3,4,5-tetrahydro-2H-benzazepin-2-yl)purine

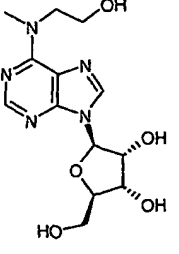
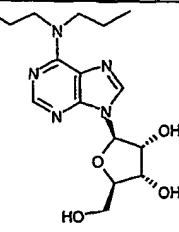
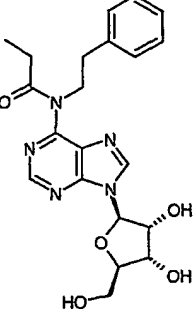
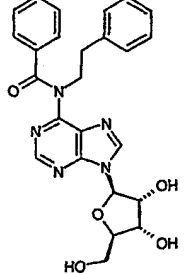
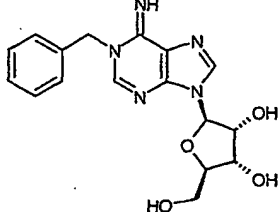
60		6-[2-(4-Cyanomethylphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine
61		6-(2,3-Dihydro-1-indolyl)-9-(β -D-ribofuranosyl)purine
62		9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl)purine
63		9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl)purine
64		6-(8-Aminosulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl)-9-(β -D-ribofuranosyl)purine

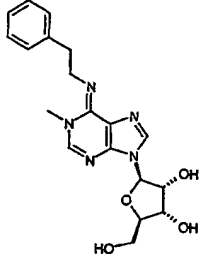
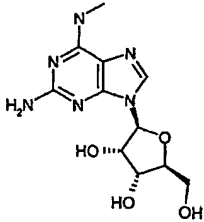
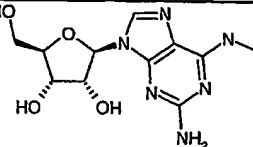
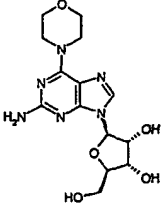
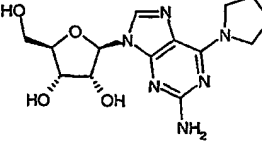
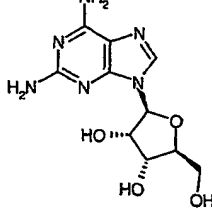
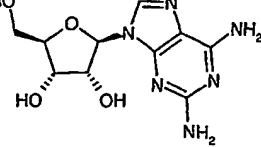
65		6-[2-(3,4-Dimethoxyphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine
66		6-[2-(4-Hydroxyphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine
67		6-(2-Isoindolinyl)-9-(β -D-ribofuranosyl)purine
68		6-(7-Aminosulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl)-9-(β -D-ribofuranosyl)purine
69		6-(N-Cyclohexylmethylamino)-9-(β -D-ribofuranosyl)purine
70		6-(N-Hexylmethylamino)-9-(β -D-ribofuranosyl)purine

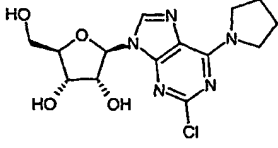
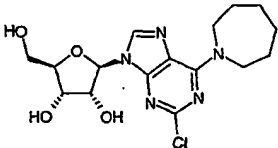
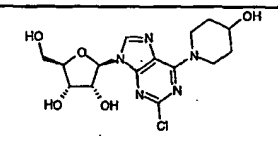
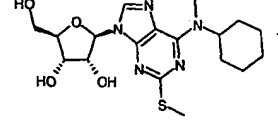
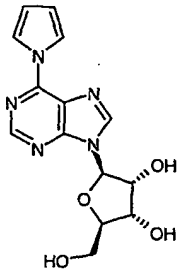
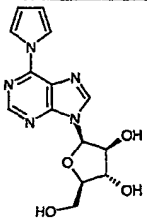
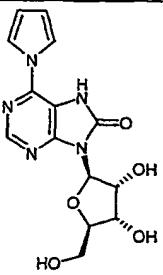
71		6-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5ylamino)-9-(β -D-ribofuranosyl)purine
72		6-[N-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5yl)methylamino]-9-(β -D-ribofuranosyl)purine
73		6-[N-(5-Aminopentyl)methylamino]-9-(β -D-ribofuranosyl)purine
74		6-[(5-Chloro-2-methoxyphenyl)methylamino]-9-(β -D-ribofuranosyl)purine
75		6-[(2-Methylphenyl)methylamino]-9-(β -D-ribofuranosyl)purine
76		6-(Hexamethyleneimino)-9-(β -D-ribofuranosyl)purine

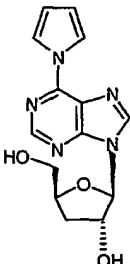
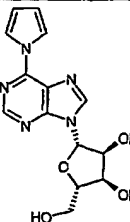
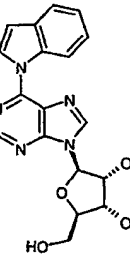
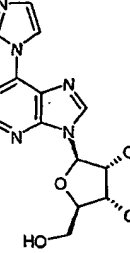
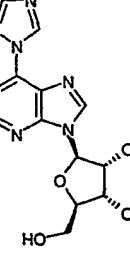
77		6-(1-Pyrrolidinyl)-9-(β -D-ribofuranosyl)purine
78		6-(4-Hydroxypiperidin-1-yl)-9-(β -D-ribofuranosyl)purine
79		6-(1-Piperidinyl)-9-(β -D-ribofuranosyl)purine
80		6-(2-Propenyl)amino-9-(β -D-ribofuranosyl)purine
81		6-(2-Propynyl)amino-9-(β -D-ribofuranosyl)purine
82		6-(1-Methyl)ethylamino-9-(β -D-ribofuranosyl)purine

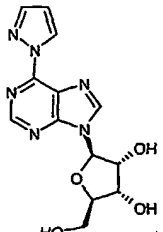
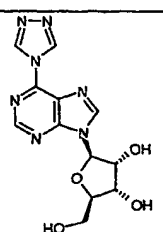
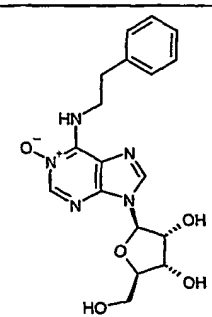
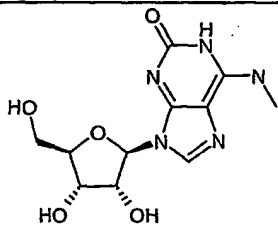
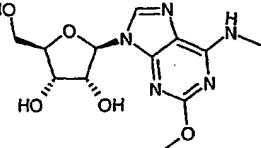
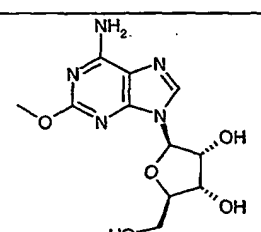
83		6-bis-(2-Propenyl)amino-9-(β -D-ribofuranosyl)purine
84		6-(2-Phenylethyl)methylamino-9-(β -D-ribofuranosyl)purine
85		6-Ethylmethylamino- 9-(β -D-ribofuranosyl)purine
86		6-bis-[(3-Methyl)butylamino]-9-(β -D-ribofuranosyl)purine
87		6-(4-Aminophenyl)methylamino-9-(β -D-ribofuranosyl)purine
88		6-(2-Pyridylmethyl)amino-9-(β -D-ribofuranosyl)purine

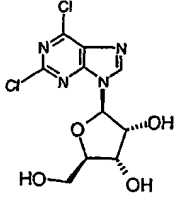
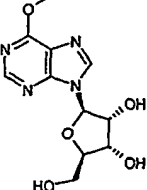
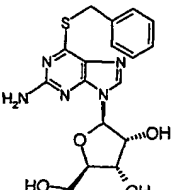
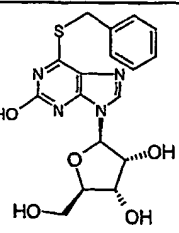
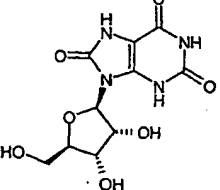
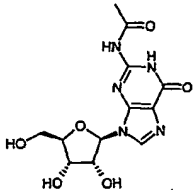
89		6-(2-Hydroxyethyl)methylamino-9-(β -D-ribofuranosyl)purine
90		6-Dipropylamino-9-(β -D-ribofuranosyl)purine
91		6-[2-Phenyl-(N-propionyl)ethylamino]-9-(β -D-ribofuranosyl)purine
92		6-(N-Benzoyl-2-phenylethylamino)-9-(β -D-ribofuranosyl)purine
93		1-Benzyl-6-imino-9-(β -D-ribofuranosyl)purine

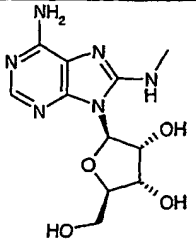
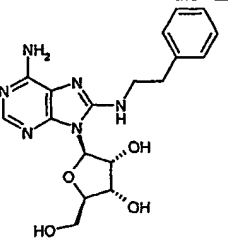
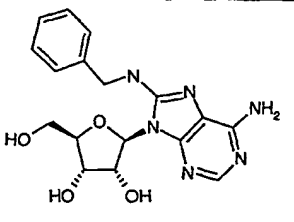
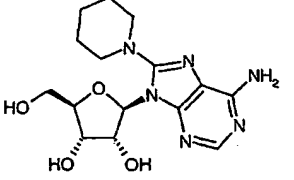
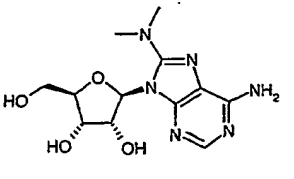
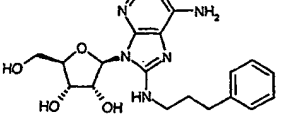
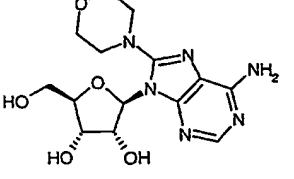
94		1-Methyl-6-(2-phenylethylimino)-9-(β -D-ribofuranosyl)purine
95		2-Amino-6-methylamino-9-(β -L-ribofuranosyl)purine
96		2-Amino-6-methylamino-9-(β -D-ribofuranosyl)purine
97		2-Amino-6-(4-morpholinyl)-9-(β -D-ribofuranosyl)purine
98		2-Amino-6-(1-pyrrolidinyl)-9-(β -D-ribofuranosyl)purine
99		2,6-Diamino-9-(β -L-ribofuranosyl)purine
100		2,6-Diamino-9-(β -D-ribofuranosyl)purine

101		2-Chloro-6-(1-pyrrolidinyl)-9-(β -D-ribofuranosyl)purine
102		2-Chloro-6-(1-hexamethyleneimino)-9-(β -D-ribofuranosyl)purine
103		2-Chloro-6-(4-hydroxy-1-piperidinyl)-9-(β -D-ribofuranosyl)purine
104		6-[(N-Cyclohexyl)methylamino]-2-methylthio-9-(β -D-ribofuranosyl)purine
105		6-(1-Pyrrolyl)-9-(β -D-ribofuranosyl)purine
106		6-(1-Pyrrolyl)-9-(β -D-arabinofuranosyl)purine
107		6-(1-Pyrrolyl)-9-(β -D-ribofuranosyl)purin-8-(7H)-one

108		9-(3-Deoxy- β -D-ribofuranosyl)-6-(1-pyrrolyl) purine
109		6-(1-Pyrrolyl)-9-(β -L-ribofuranosyl)purine
110		6-(1-Indolyl)-9-(β -D-ribofuranosyl)purine
111		6-(1-Imidazolyl)-9-(β -D-ribofuranosyl)purine
112		9-(β -D-Ribofuranosyl)-6-(1,2,4-triazol-1-yl)purine

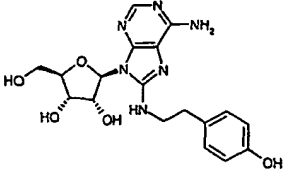
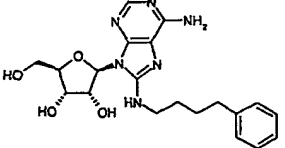
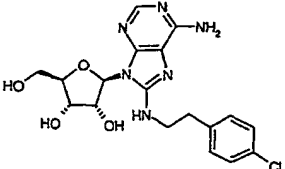
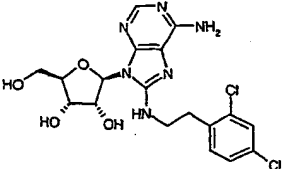
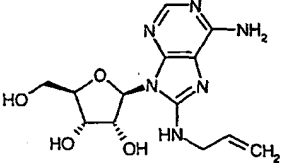
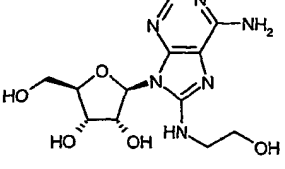
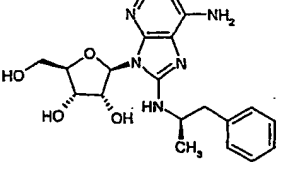
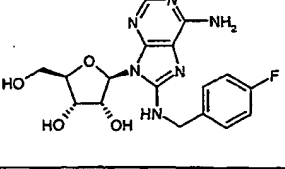
113		6-(1-Pyrazolyl)- 9-(β -D-ribofuranosyl)purine
114		9-(β -D-ribofuranosyl) 6-(1,2,4-triazol-4-yl)purine
115		6-(2-Phenylethylamino)- 9-(β -D-ribofuranosyl)purine-1-oxide
116		6-Methylamino-9-(β -D-ribofuranosyl)purin-2(1H)-one
117		2-Methoxy-6-methylamino-9-(β -D-ribofuranosyl)purine
118		2-Methoxyadenosine

119		2,6-Dichloro-9-(β-D-ribofuranosyl)purine
120		6-Methoxy-9-(β-D-ribofuranosyl)purine
121		2-Amino-6-benzylthio-9-(β-D-ribofuranosyl)purine
122		6-Benzylthio-2-hydroxy-9-(β-D-ribofuranosyl)purine
123		9-(β-D-Ribofuranosyl)purine-2,6,8(1H,3H,7H)-trione
124		2-(Acetylamino)inosine

125		8-(Methylamino)adenosine
126		8-(2-Phenylethylamino)adenosine
127		8-Benzylaminoadenosine
128		8-(1-Piperidinyl)adenosine
129		8-(Dimethylamino)adenosine
130		8-(3-Phenylpropylamino)adenosine
131		8-(4-Morpholinyl)adenosine

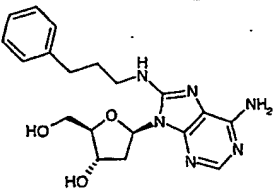
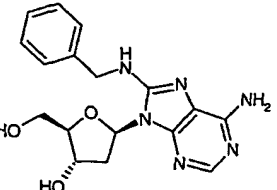
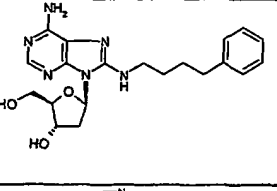
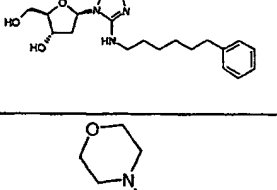
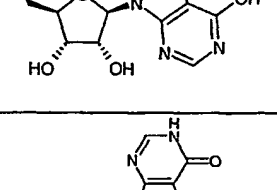
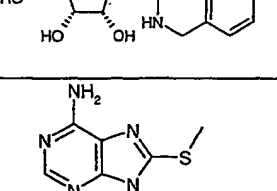
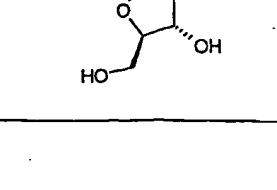
132		8-(N-Methyl-2-phenylethylamino)adenosine
133		8-(3-Pyridylmethylamino)adenosine
134		8-(Ethylamino)adenosine
135		8-(1,2,3,4-Tetrahydro-2-isoquinolyl)adenosine
136		8-[2-(4-Morpholinyl)ethylamino]adenosine
137		8-(Hexylamino)adenosine
138		8-(2-Cyclohexylethylamino)adenosine

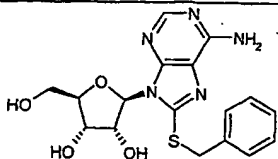
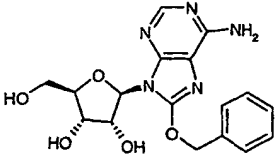
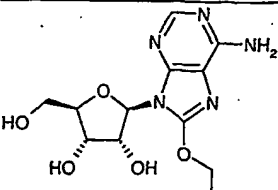
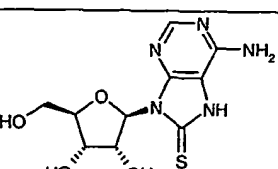
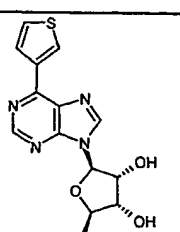
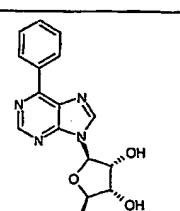
139		8-(2(R,S)-Phenylpropylamino) adenosine
140		8-[2-(4-Methylphenyl) ethylamino]adenosine
141		8-[2-(1-Methyl-2-pyrrolyl) ethylamino]adenosine
142		8-[2-(4-Aminosulphonylphenyl) ethylamino]adenosine
143		8-(4-Phenyl-1-piperazinyl) adenosine
144		8-(2-(4-Imidazolyl)adenosine
145		8-(1-Naphthylmethylamino)adenosine

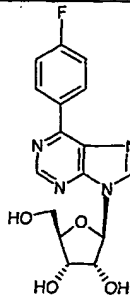
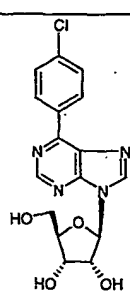
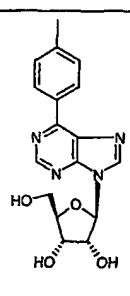
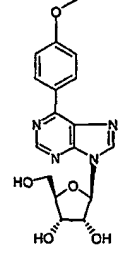
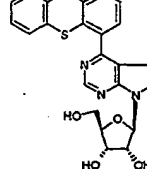
146		8-[2-(4-Hydroxyphenyl)ethylamino]adenosine
147		8-(4-Phenylbutylamino)adenosine
148		8-[2-(4-Chlorophenyl)ethylamino]adenosine
149		8-[2-(2,4-Dichlorophenyl)ethylamino]adenosine
150		8-(2-Propenylamino)adenosine
151		8-(2-Hydroxyethylamino)adenosine
152		8-(1(R)-Methyl-2-phenylethylamino)adenosine
153		8-(4-Fluorobenzylamino)adenosine

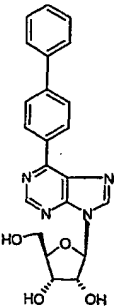
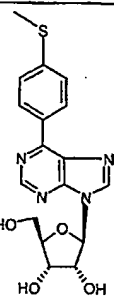
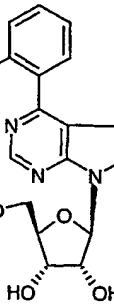
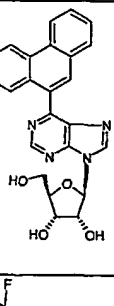
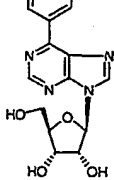
154		8-[(4-Hydroxycarbonyl)benzylamino]adenosine
155		8-(2-Propynylamino)adenosine
156		8-(1-Methylethylamino)adenosine
157		8-[(4-Trifluoromethyl)benzylamino]adenosine
158		8-[(2,5-Dimethoxy)benzylamino]adenosine
159		8-[2-(2-thienyl)ethylamino]adenosine
160		8-[2-(4-Aminophenyl)ethylamino]adenosine

161		8-(2-Phenoxyethylamino)adenosine
162		8-[(2-Thienyl)methylamino)adenosine
163		8-[(4-tert-Butyl)benzylamino]adenosine
164		8-(1(R)-Phenylethylamino)adenosine
165		8-(1(S)-Phenylethylamino)adenosine
166		8-(6-Phenylhexylamino)adenosine
167		8-[2-Hydroxy-1(S)-phenyl]ethylamino)adenosine
168		2'-Deoxy-8-(2-phenylethylamino)adenosine

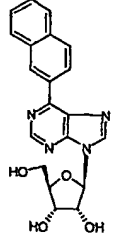
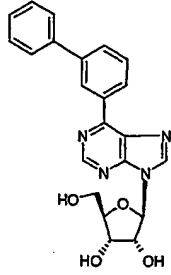
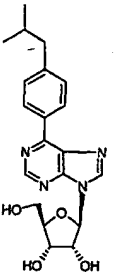
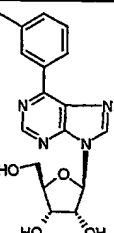
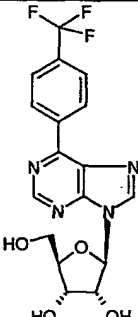
169		2'-Deoxy-8-(3-phenylpropylamino)adenosine
170		8-Benzylamino-2'-deoxyadenosine
171		2'-Deoxy-8-(4-phenylbutylamino)adenosine
172		2'-Deoxy-8-(6-phenylhexylamino)adenosine
173		8-(4-Morpholinyl)inosine
174		8-(Benzylamino)inosine
175		8-(Methylthio)adenosine

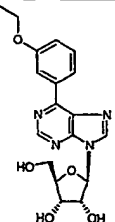
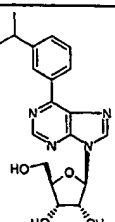
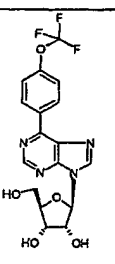
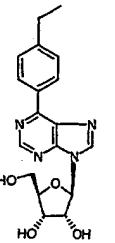
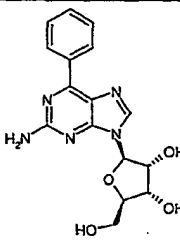
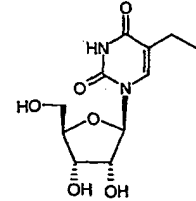
176		8-(Benzylthio)adenosine
177		8-(Benzylxy)adenosine
178		8-Ethoxyadenosine
179		6-Amino-9-(β -D-ribofuranosyl)purine-8(7H)-thione
180		8-[(1-Hydroxy-1-methyl)ethyl]adenosine
181		9-(β -D-Ribofuranosyl)-6-(3-thienyl)purine
182		6-Phenyl-9-(β -D-ribofuranosyl)purine

183		6-(4-Fluorophenyl)-9-(β -D-ribofuranosyl)purine
184		6-(4-Chlorophenyl)-9-(β -D-ribofuranosyl)purine
185		6-(4-Methylphenyl)-9-(β -D-ribofuranosyl)purine
186		6-(4-Methoxyphenyl)-9-(β -D-ribofuranosyl)purine
187		9-(β -D-Ribofuranosyl)-6-(1-thianthrenyl)purine

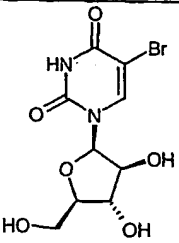
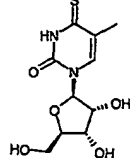
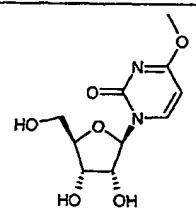
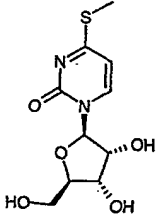
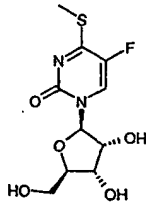
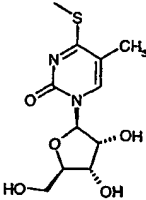
188		6-(4-Biphenyl)-9-(β -D-ribofuranosyl)purine
189		6-(4-Methylthiophenyl)-9-(β -D-ribofuranosyl)purine
190		6-(2-Methylphenyl)-9-(β -D-ribofuranosyl)purine
191		6-(9-Phenanthrenyl)-9-(β -D-ribofuranosyl)purine
192		9-(β -D-Ribofuranosyl)-6-(3-trifluoromethylphenyl)purine

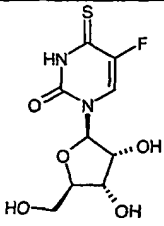
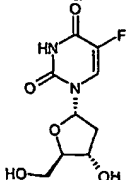
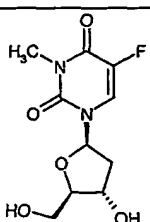
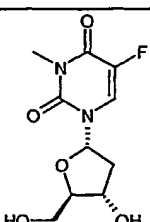
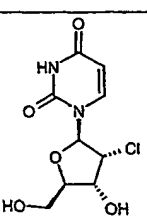
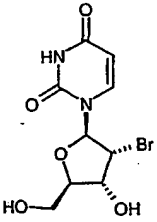
193		6-(2-Phenoxyphenyl)-9-(β-D-ribofuranosyl)purine
194		6-(4-tert-Butylphenyl)-9-(β-D-ribofuranosyl)purine
195		9-(β-D-Ribofuranosyl)-6-(2-trifluoromethoxyphenyl)purine
196		6-(4-Phenoxyphenyl)-9-(β-D-ribofuranosyl)purine
197		6-(3-Methoxyphenyl)-9-(β-D-ribofuranosyl)purine

198		6-(2-Naphthyl)-9-(β -D-ribofuranosyl)purine
199		6-(3-Biphenylyl)-9-(β -D-ribofuranosyl)purine
200		6-[4-(2-Methylpropyl)phenyl]-9-(β -D-ribofuranosyl)purine
201		6-(3-Fluorophenyl)-9-(β -D-ribofuranosyl)purine
202		9-(β -D-Ribofuranosyl)-6-(4-trifluoromethylphenyl)purine

203		6-(3-Ethoxyphenyl)-9-(β -D-ribofuranosyl)purine
204		6-[3-(1-Methyl)ethylphenyl]-9-(β -D-ribofuranosyl)purine
205		9-(β -D-Ribofuranosyl)-6-(4-trifluoromethoxyphenyl)purine
206		6-(4-Ethylphenyl)-9-(β -D-ribofuranosyl)purine
207		2-Amino-6-phenyl-9-(β -D-ribofuranosyl)purine
208		5-Ethyluridine

209		5-[(1-Methyl)ethyl]uridine
210		5-Methoxymethyluridine
211		5-Ethoxymethyluridine
212		5-Chlorouridine
213		5-Methyl-1-(β-L-ribofuranosyl)uracil
214		1-(β-D-Arabinofuranosyl)-5-ethyluracil

215		1- (β -D-Arabinofuranosyl)-5-bromouracil
216		5-Methyl-4-thiouridine
217		4-Methoxy-1- (β -D-ribofuranosyl)pyrimidin-2(1H)-one
218		4-Methylthio-1- (β -D-ribofuranosyl)pyrimidin-2(1H)-one
219		5-Fluoro-4-methylthio-1- (β -D-ribofuranosyl)pyrimidin-2(1H)-one
220		5-Methyl-4-methylthio-1- (β -D-ribofuranosyl)pyrimidin-2(1H)-one

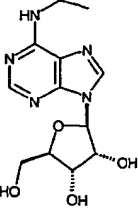
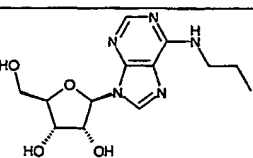
221		5-Fluoro-4-thiouridine
222		1-(2-Deoxy- α -D-erythro-pentofuranosyl)-5-fluorouracil
223		2'-Deoxy-5-fluoro-3-methyluridine
224		1-(α -D-Erythro-2-deoxypentofuranosyl)-5-fluoro-3-methyluracil
225		2'-Chloro-2'-deoxyuridine
226		2'-Bromo-2'-deoxyuridine

227		1-(2-Deoxy- β -D-lyxofuranosyl)-5-methyluracil
228		3'-Deoxy-3'-fluoro-5-methyluridine
229		2',3'-Dideoxy-5-ethyl-3'-methoxyuridine
230		5'-Benzylxy-2',3'-dideoxy-5-methyluridine
231		2',3'-Dideoxy-5-ethyl-3'-iodouridine
232		3'-Azido-2',3'-dideoxy-5-ethyluridine

233		3'-Azido-2',3'-dideoxy-5-methylcytidine
234		1-(3-Deoxy- β -L-threo-pentofuranosyl)-5-fluorocytosine
235		4-Methylamino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
236		5-Fluoro-4-methylamino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
237		4-(1-Pyrrolyl)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
238		4-Oximino-1-(β -L-ribofuranosyl)pyrimidin-2(1H)-one

239		4-Oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
240		4-Oximino-1-(β -D-arabinofuranosyl)pyrimidin-2(1H)-one
241		5-Fluoro-4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
242		1-(2-Deoxy-2,2-difluoro- α -D-erythropentofuranosyl)uracil
243		1-(2-Deoxy-2,2-difluoro- β -D-erythropentofuranosyl)cytosine
244		L-Cytidine

245		4-Amino-1-(2,2-difluoro-3-hydroxy-4-hydroxymethylcyclopentyl)-1H-pyrimidin-2-one
246		4-Amino-1(R)-(2(S),3(R)-dihydroxy-4(R)-hydroxymethylcyclopentyl)-1H-pyrimidin-2-one
247		1-(β-D-Xylofuranosyl)cytosine
248		1-(3-Deoxy-3-fluoro-β-D-xylofuranosyl)uracil
249		1-(3-Deoxy-3-fluoro-β-D-xylofuranosyl)cytosine
250		3'-Deoxy-3'-hydroxymethylcytidine
251		2'-Deoxy-2'-methoxyuridine

252		6-Ethylamino-9-(β -D-ribofuranosyl)purine
253		6-Propylamino-9-(β -D-ribofuranosyl)purine

The compounds of formula I according to the present invention are prepared as follows:

5 The compounds of formula I may be prepared by various methods known in the art of organic chemistry in general and nucleoside analogue synthesis in particular. The starting materials for the syntheses are either readily available from commercial sources or are known or may themselves be prepared by techniques known in the art. General reviews of the preparation of nucleoside analogues are included in the following:

10 A M Michelson "The Chemistry of Nucleosides and Nucleotides", Academic Press, New York 1963.

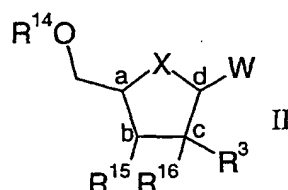
L Goodman "Basic Principles in Nucleic Acid Chemistry" ed P O P Ts'0, Academic Press, New York 1974, Vol. 1, chapter 2.

"Synthetic Procedures in Nucleic acid Chemistry" ed W W Zorbach and R S Tipson, Wiley, New York, 1973, Vol. 1 and 2.

15 The synthesis of carbocyclic nucleosides has been reviewed by: L Agrofoglio et al Tetrahedron, 1994, 50, 10611.

The strategies available for the synthesis of compounds of formula I include:

1. Condensation of a protected furanose, thiofuranose or cyclopentane derivative of formula II



20

wherein

R³ is as defined above;

R¹⁴ is a hydroxy protecting group;

25 R¹⁵ is as defined for R¹ except that when R¹ is hydroxy R¹⁵ is a group OR¹⁷ wherein R¹⁷ is a hydroxy protecting group;

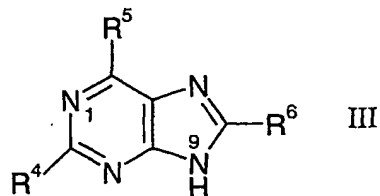
R^{16} is as defined for R^2 except that when R^2 is hydroxy R^{16} is a group OR^{17} wherein R^{17} is a hydroxy protecting group;

X is O, S or CH_2 ;

5 W is a leaving group such as acyloxy, aryloxy, alkylsulphonate, arylsulphonate, S-benzyl or halogen; and

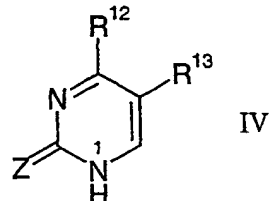
a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents;

with an appropriate purine of formula III



10 wherein R^4 , R^5 and R^6 are as defined in formula I;

or pyrimidine of formula IV



wherein Z , R^{12} and R^{13} are as defined in formula I;

15 or a derivative of the purine or pyrimidine such as for example a heavy metal or silyl derivative.

20 The particular nature of the hydroxy protecting groups R^{14} or R^{17} is selected in accordance with conventional techniques. Examples for hydroxy protecting groups are acyl (e.g. acetyl), aroyl (e.g. benzoyl), ether (e.g. bis-acetonide), silylether (e.g. trimethylsilyl, tert-butyldimethylsilyl) or arylmethyl (e.g. benzyl, triphenylmethyl).

5 The condensation reaction may be performed using standard methods including the use of a Lewis acid catalyst such as mercuric bromide or stannic chloride or trimethylsilyltrifluoromethane sulphonate in solvents such as acetonitrile, 1,2-dichloroethane, dichloromethane, chloroform or toluene at reduced, ambient or elevated temperature. Examples for the condensation reaction of a protected furanose or thiofuranose of formula II where X is O or S with an appropriate pyrimidine or purine derivative are as follows:

- 10 a) The reaction may be performed by the condensation of heavy metal derivatives of purines of formula III or pyrimidines of formula IV (e.g. chloromercuri derivatives) with a compound of formula II as described by J Davoll and B A Lowry J Am Chem Soc 1951, 73, 1650; J J Fox, N Yung, J Davoll and G B Brown J Am Chem Soc 1956, 78, 2117.
- 15 b) The reaction may also involve the condensation of alkoxy pyrimidines with compounds of formula II as described by K A Watanabe, D H Hollenberg and J J Fox Carbohydrates, Nucleosides and Nucleotides 1974, 1,1.
- 20 c) The reaction may be performed by the condensation of silyl derivatives of purines of formula III or pyrimidines of formula IV with compounds of formula II as described by U Niedballa and H Vorbruggen J Org Chem 1976, 41, 2084; U Niedballa and H Vorbruggen J Org Chem 1974, 39, 3672. A J Hubbard, A S Jones and R T Walker Nucleic Acids Res 1984, 12, 6827.
- 25 d) Fusion of per-acylated sugars with purines under vacuum in the presence of p-toluene sulphonic acid has been described by T Simadate, Y Ishudo and T Sato Chem Abs 1962, 56, 11 692 and W Pfleiderer, R K Robins Chem Ber 1965, 98, 1511.
- e) Further coupling reactions have been described by K A Watanabe, D H Hollenberg and J J Fox Carbohydrates, Nucleosides and Nucleotides 1974, 1,1.

 Examples for the condensation reaction of a protected cyclopentane derivative of formula II wherein X is CH₂ with an appropriate purine derivative of formula III or pyrimidine derivative of formula IV are as follows:

- 30 a) The nucleophilic displacement of the leaving group W in a compound of formula II where X is CH₂ with a purine derivative of formula III or pyrimidine

derivative of formula IV as described by H Kapeller, H Baumgartner and H Griengl, Monatsh Chem, 1997, 128, 191 and P Wang et al, Tet Lett 1997, 38, 4207.

5 b) The reaction of a cyclopentane derivative of formula II in which W is OH with a purine derivative under Mitsonobu conditions, which employs a triarylphosphine such as triphenyl phosphine and a diazodicarboxylic acid diester such as diethyl azodicarboxylate as reagents, as described by T Jenny et al Helv Chim Acta 1992, 25, 1944.

10 Such methods often result in mixtures of anomeric nucleoside derivatives which can be separated by standard techniques known to the art such as recrystallisation, column chromatography, high performance liquid chromatography or super critical fluid chromatography.

15 The purine derivatives of formula III and pyrimidines derivatives of formula IV for above condensation reactions can be obtained commercially or can be prepared by procedures known to the art.

The preparation of purine derivatives of formula III is reviewed by G Shaw in "Comprehensive Heterocyclic Chemistry" pub Pergamon Press Vol. 5 chapter 4.09, p 499 and "Comprehensive Heterocyclic Chemistry II" pub Pergamon Press Vol 7, chapter 7.11 p 397.

20 The preparation of pyrimidines derivatives of formula IV is reviewed by D J Brown "The Chemistry of Heterocyclic Compounds – The Pyrimidines" 1962 and Supplement 1, 1970, pub John Wiley and Sons, New York, by D J Brown in "Comprehensive Heterocyclic Chemistry" pub Pergamon Press Vol. 5 chapter 4.09, p 499 and by K Unheim and T Benneche in "Comprehensive Heterocyclic Chemistry II" pub Pergamon Press Vol. 6 chapter 6.02 p 93.

25 For example the appropriate purine base of formula III may be prepared from the corresponding purine wherein the 2, 6 or 8 position of the purine base is substituted with a suitable leaving group such as halogen or sulphonate. Such purine precursors bearing leaving groups are available commercially e.g. 6-chloropurine (Aldrich Chemical Company), 2,6-dichloropurine (Aldrich Chemical Company), 2-chloro-6-aminopurine (Aldrich Chemical Company), 8-bromoadenine (Sigma-Aldrich Company Limited) or obtained by procedures

known in the art. For example 2- and 6-chloro substituted purines can be prepared by chlorination of the corresponding 2 and 6-hydroxypurines respectively by the use of chlorinating agents such as phosphorus oxychloride (D S Bakuni et al Indian J Chem Sect B 1984, 23, 1286; M P LaMontagne et al J Heterocycl Chem 1983, 20, 295) while introduction of a bromine into the 8-position of purines can be accomplished by direct bromination using brominating agents such as for example bromine (M Mano et al, Chem Pharm Bull 1983, 31, 3454) or N-bromosuccinimide (J L Kelley et al J Heterocycl Chem 1990, 27, 1505). The purines where the 6 substituent is alkoxy, aryloxy, SH, alkylthio, arylthio, alkylamino, cycloalkylamino, saturated cyclic amino, nitrogen linked heteroaromatic, hydroxylamino, alkoxyamino, hydrazine, alkylhydrazino may be prepared by treatment of the corresponding 6-halopurine with the appropriate alkoxides, thiols, amines, nitrogen containing heterocycles, hydroxylamines and hydrazines, (e.g M-Y Chae et al J Med Chem, 1994, 37, 342; G Niebch and F Schneider, Z.Naturforsch. B. Anorg. Chem. Org. Chem. Biochem. Biophys. Biol. 1972, 27, 675; M P LaMontagne et al, J Heterocycl Chem 1983, 20, 295; K G Estep et al J Med Chem 1995, 38, 2582). Similarly 2-substituted purines can be prepared from the corresponding 2-halopurine for example purines where the 2 substituent is alkoxy, aryloxy, SH, alkylthio, arylthio or NR⁷R⁸ can be prepared from the corresponding 2-halopurine by treatment with alkoxides, thiols or amines (e.g. G B Barlin and D M Fenn, Aust J Chem, 1983, 36, 633; D A Nugiel et al, J Org Chem, 1997, 62, 201). Similarly 8-substituted purines can be prepared from the corresponding 8-halopurine. For example purines where the 8-substituent is alkoxy, aryloxy, SH, alkylthio, arylthio or NR⁷R⁸ can be prepared by treatment of the corresponding 8-bromopurine with the appropriate alkoxides, thiols or amines (Xing et al, Tet Lett, 1990, 31, 5849; M Mano et al, Chem Pharm Bull 1983, 31, 3454). Where the 2, 6 or 8 substituent is a cyclic amine moiety the purine can be prepared from the 6-aminopurine by reaction with an appropriate dialkylating agent such as a dihaloalkane. In some cases where the 6-substituent is a nitrogen containing heteroaromatic linked through the nitrogen atom the purine may be prepared from the 6-aminopurine by reaction with a dicarbonyl compound or a reactive derivative of this such as an acetal. For example 6-(1H-pyrrol-1-yl)-1H-purine can be prepared from 6-chloropurine by reaction with 2,5-dimethoxytetrahydrofuran as described by K G Estep et al J Med Chem 1995, 38, 2582.

The furanose and thiofuranose derivatives of formula II used for the condensation reactions can be prepared by methods known in the art of carbohydrate chemistry.

5 Furanose derivatives can be prepared from commercially available carbohydrate starting materials such as the D or L forms of ribose, arabinose, xylose or lyxose. Following introduction of protecting groups which are compatible with the chemistry, modification of either the 2-hydroxy substituent or 3-hydroxy substituent is possible. For example direct alkylation with alkylating agents such as alkyl halides, alkyl sulphonates or diazoalkanes provides the corresponding O-alkyl derivatives as exemplified by M E Jung, C Castro, S I Khan, Nucleosides and Nucleotides; 1998, 17, 2383; G Parmentier, G Scmitt, F Dolle, B Luu Tet 1994, 50, 5361. Conversion of either hydroxy to a leaving group such as halo followed by reduction provides the 2- or 3-deoxysugar derivatives as described by K C Nicolaou et al J Am Chem Soc 1988, 110, 4672. Also conversion of either hydroxy to a leaving group such as halo or sulphonate by standard methods followed by displacement with nucleophilic reagents for example sodium or lithium azide to introduce an azido group (A M Ozols et al, Synthesis, 1980, 557). Direct introduction of a fluorine substituent can be accomplished with fluorinating agents such as diethylaminosulphur trifluoride as described by F Puech, G Gosselin and J-L Imbach Tet Lett 1989, 30, 3171 or conversion of the hydroxy substituent to a leaving group such as halo or sulphonate and displacement using reagents such as tetrabutylammonium fluoride as described in Tet Asym 1990, 1 715.

10

15

20

25 3'-Alkyl substituted furanoses can be prepared by construction of the sugar ring from γ -hydroxymethyl- γ -butyrolactone as described by K Ayei-Aye and D C Baker, Carbohydr Res 1988, 183, 261 and by M Okabe et al J Org chem, 1988, 53, 4780. Alternatively, cyclohexenecarboxylic acid derivatives can be used as described by K C Schneider and S A Benner, Tet Lett, 1990, 31, 335.

30 3'-hydroxymethyl substituted furanoses can be synthesised from 3-[(4-bromobenzyl)oxy]methyl]oxirane-2-methanol as described by L Svansson et al, J Org Chem 1991, 56, 2993.

2,2-Difluorofuranose derivatives can be prepared from D-glucose or D-mannose as described by R Fernandez, M I Mateu, R Echarri and S Castillon Tet 1998, 54, 3523. The thiofuranose derivatives of formula II where X is S can be

prepared by literature procedures such as L Bellon, J L Barascut, J L Imbach Nucleosides and Nucleotides 1992, 11, 1467 and modified in a similar fashion to the furanose analogues described above.

5 The cyclopentane derivatives of formula II where X is CH₂ can be prepared by methods known in the art of organic chemistry and by methods and references included in L Agrofolio et al Tetrahedron 1994, 50, 10611.

2. Construction of the heterocyclic base after glycosylation.

Such methods include :

- 10 a) those which for example utilise furanosylamine derivatives as described by N J Cusack, B J Hildick , D H Robinson, P W Rugg and G Shaw JCS Perkin I 1973, 1720 or G Shaw, R N Warrener, M H Maguire and R K Ralph, J Chem Soc 1958, 2294.
- 15 b) those which utilise for example furanosylureas for pyrimidine nucleoside synthesis as described by J Šmejkal, J Farkas, and F Šorm Coll Czech Chem Comm 1966, 31, 291.
- c) The preparation of purine nucleosides from imidazole nucleosides as reviewed by L B Townsend Chem Rev 1967, 67, 533.
- 20 d) the preparation of compounds of formula I wherein X is CH₂ can be accomplished from 1-hydroxymethyl-4-aminocyclopentane derivatives as described by Y F Shealy and J D Clayton J Amer Chem Soc 1969, 91, 3075; R Vince and S Daluge J Org Chem 1980, 45, 531; R C Cermak and R Vince Tet Lett 1981,2331; R D Elliott et al J Med Chem 1994,37, 739; A D Borthwick et al, J Med Chem 1990, 33, 179.

25

3. Modification or inter-conversion of preformed nucleosides.

A. Modification of the purine or pyrimidine base moiety.

Methods include:

- 5 a) the deamination of aminopurine or aminopyrimidine nucleosides as described by J R Tittensor and R T Walker European Polymer J 1968, 4, 39 and H Hayatsu Progress in Nucleic Acid Research and Molecular Biology 1976, Vol. 16, p75.
- 10 b) The conversion of the 4-hydroxy group of 4-hydroxypyrimidine nucleosides to a leaving group and displacement with nucleophilic reagents. Such leaving groups include halogen as described by J Brokes and J Beranek Col Czech Chem Comm 1974, 39, 3100 or 1,2,4-triazole as described by K J Divakar and C B Reece J Chem Soc Perkin Trans I 1982, 1171.
- 15 c) 5-substitution of pyrimidine nucleosides has been achieved by the use of 5-metalloc derivatives such as 5-mercuri or 5-palladium for example as described by D E Bergstrom and J L Ruth J Amer Chem Soc 1976, 98, 1587. Introduction of fluoro into the 5 position of pyrimidine nucleosides can be achieved with reagents such as trifluoromethyl hypofluorite as described by M J Robins Ann New York Acad Sci 1975, 255, 104.
- 20 d) modified purine nucleosides may be prepared from the corresponding purine nucleoside derivatives wherein the 2, 6 or 8 substituent is a suitable leaving group such as halogen or sulphonate or 1,3,4-triazole. Thus the compounds for example where the purine 6 substituent is alkoxy, aryloxy, SH, alkylthio, arylthio, alkylamino, cycloalkylamino hydroxylamino, alkoxyamino or hydrazino may be prepared by treatment of the appropriate 6-halopurine or 6-(1,2,4-triazol-4-yl)purine nucleoside derivatives with the appropriate alcohols, thiols or amines, hydroxylamines or hydrazines. Such conversions are described by V Nair and A J Fassbender Tet 1993, 49, 2169 and by V Samano, R W Miles and M J Robins J Am Chem Soc 1994, 116, 9331. Where the 6 substituent is a cyclic amine or aromatic amine moiety the purine nucleoside analogue can be prepared from the 6-aminopurine nucleoside derivative by reaction respectively with an appropriate dialkylating agent such as a dihaloalkane or with a dicarbonyl compound or a reactive derivative of this such as an acetal. For example as described by M Haidoune and R Mornet J Heterocyclic Chem 1995, 31, 1462. Similarly 8-substituted purine nucleosides can be prepared by treatment of the corresponding 8-
- 25
- 30
- 35

- halopurine nucleoside with the appropriate nucleophilic reagent for example alkoxides, thiols or amines as described by L Tai-Shun, C Jia-Chong, I Kimiko and A C Sartorelli J Med Chem 1985, 28, 1481; Nandanam et al J Med Chem 1999, 42, 1625; J Jansons, Y Maurinsh, and M Lidaks Nucleosides and Nucleotides 1995, 14, 1709. Introduction of a 8-cyano substituent can be accomplished by displacement of using a metal cyanide as described by L-L Gundersen, Acat Chem Scand 1996, 50, 58. 2-modified purine nucleoside may be prepared in a similar fashion as described by T Steinbrecher, C Wamelung, F Oesch and A Seidl Angew Chem Int Ed Engl 1993, 32, 404.
- 10
- e) Where the substituent at the 2, 6 or 8-position of the purine nucleoside is linked via a carbon carbon bond e.g. alkyl or aryl then metal catalysed cross-coupling procedures can be used starting with the appropriate 2, 6 or 8-halosubstituted purine nucleoside analogue. Such procedures are described by AA Van Aerschott, et al J Med Chem 1993, 36, 2938; D E Bergstrom and P A Reday Tet Lett 1982, 23, 4191. M Hocek, A Holy, I Votruba and H Dvarakova J Med Chem 2000, 43, 1817. C Tu, C Keane and B E Eaton Nucleosides and Nucleotides 1995, 14, 1631.
- 15
- f) Oxidation of the 3-nitrogen in pyrimidine nucleoside analogues or 1-nitrogen in purine nucleoside derivatives can be accomplished using hydrogen peroxide or organic peroxides as described by G B Brown Progress in Nucleic Acid Research and Molecular Biology ed J N Davidson and W E Cohn, Academic Press, New York 1968, 8, 209.
- 20
- g) Alkylation of the 3-nitrogen in uracil nucleoside analogues can be accomplished using alkylating agents such as diazoalkanes (Miles, Biochim Biophys Acta, 1956, 22, 247), alkyl sulphonates (Scannel et al, Biochim Biophys Acta, 1959, 32, 406) or alkyl halides (Anderson et al J Chem Soc 1952, 369). Alkylation of the 3-nitrogen in cytosine nucleoside analogues can similarly be accomplished using alkylating agents such as trialkyl sulphonium halides (K Yamauchi, J Chem Soc Perkin Trans 1, 1980, 2787) or epoxides (W Zhan et al Chem Res Toxicol, 1998, 8, 148). Similarly alkylation of purine nucleoside analogues on the 1-nitrogen can be accomplished using alkylating agents such as alkyl halides (W A Szarek et al Can J Chem 1985, 63, 2149) or alkyl sulphonates (M Kawana et al J Chem Soc Perkin Trans 1, 1992, 4, 469). Aryl substituents can be
- 25
- 30
- 35

introduced onto the 1-nitrogen of purine nucleosides or the 3-nitrogen of pyrimidine nucleosides by direct arylation using aryl halides in the presence of a copper catalyst such as copper(I) oxide as described for example by T Maruyama et al, Nucleosides and Nucleotides, 1997, 16, 1079 and by T Maruyama et al J Chem Soc Perkin Trans I, 1995, 733.

4. B. Modification of the carbohydrate moiety.

Methods include:

- a) Following introduction of protecting groups which are compatible with the further chemistry, modification of either the 2'-hydroxy substituent or 3'-hydroxy substituent in the nucleoside analogue is possible. For example direct alkylation with alkylating agents such as alkyl halides, alkyl sulphonates or diazoalkanes provides the corresponding O-alkyl derivatives as exemplified by C G Edmonds et al J Chem Soc Chem Comm 1987, 12, 909; P J L M Quaedfiege et al J Org Chem 1991, 56, 5846. Conversion of either hydroxy to a leaving group such as halo by reaction with for example triphenyl phosphine and a tetrahaloalkane as described for example by L De Napoli et al, Nucleosides and Nucleotides, 1993, 12, 981, followed by reduction provides the 2- or 3-deoxysugar derivatives as described by D G Norman and C B Reese, Synthesis 1983, 304.
 - 10
 - 15
 - 20
 - 25
 - 30
 - 35
- Alternatively derivatisation of the hydroxy function by conversion to a thiocarbonate group such as phenoxy thiocarbonate or imidazoylthiocarbonate followed by reduction using free radical reducing agents such as trialkyltin hydrides as described by D H R Barton and R Subranian J Chem Soc Chem Comm 1976, 867. Direct introduction of a fluorine substituent can be accomplished with fluorinating agents such as diethylaminosulphur trifluoride as described by P Herdewijin, A Van Aerschot and L Kerremans Nucleosides and Nucleotides 1989, 8, 65. Conversion of the hydroxy substituent to a leaving group such as halo or sulphonate also allows displacement using nucleophilic reagents such as tetrabutylammonium fluoride, lithium azide, tert butyl isocyanide or metal cyanides as exemplified by H Hrebabecky, A Holy and e de Clercq Collect Czech Chem Comm 1990, 55, 1800; K E B Parkes and K Taylor Tet Lett 1988, 29, 2995. Such nucleophilic reactions can also be carried out on 2',3'-epoxynucleosides as exemplified by Huang et al J Med Chem

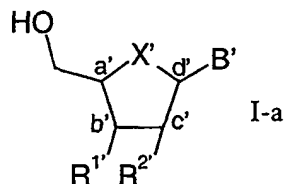
1991, 34, 1640 or using 2,3'-anhydropurimidine nucleosides as typified by Colla et al Eur J Med Chem Chim Ther 1985, 20, 295.

- 5 b) Following introduction of appropriate protecting groups on the 3' and 5'-hydroxy groups of a preformed nucleoside it is possible to oxidise the unprotected 2'-hydroxy group to a ketone using methods similar to those described by F Hansske, M D Fritz and M J Robins, Tetrahedron 1984, 40, 125. Reaction of the resultant 2'-keto nucleoside with olefination reagents such as methyl triphenyl phosphonium bromide in the manner of S Czernecki, L Mulard, J-M Valery, and A Commercon, Can.J.Chem 10 1993, 71, 413 provides the 2'-deoxy-2'-methylidene nucleoside derivatives.
- 15 c) Reaction of 2'-keto nucleosides with fluorinating agents such as diethylamino sulfur trifluoride can be used to prepare 2',2'-difluoronucleosides as described by D Bergstrom, E Romo and P Shum Nucleosides and Nucleotides 1987, 6,53.
- 20 d) The principal methods of introducing an alkyl group into the 3'-position of nucleosides involve, free-radical coupling of protected nucleosides which are suitably derivatised in the 3'-position, for example from 3'-iodonucleosides as described by D Yu and M d'Alarco, J Org Chem 1989, 54,3240 or from 3'-O-phenoxythiocarbonyl nucleosides as described by J Fiandor and S Y Tam, Tet Lett, 1990,31, 597 and C K Chu et al, J Org Chem, 1989,54, 2767, or through addition of cyanide to 3'-ketonucleosides as described by M J Camarasa et al, J Med Chem, 1989, 32, 1732. A 3'-hydroxymethyl substituent can be introduced by reduction of the corresponding 3'-C-formyl nucleoside as described by M J Bamford et al, J Med Chem, 1990, 33, 2494. The 3'-C-formyl nucleoside can be produced in turn by elaboration of 3'-keto nucleosides or from 2',3'-anhydronucleosides.
- 30 The preformed nucleoside derivatives are either available commercially or synthesised in accordance with the methods described above.

Also part of this invention are novel purine and pyrimidine nucleoside derivatives, a process for their manufacture, pharmaceutical compositions and the use of such compounds in medicine. In particular, the compounds are useful as inhibitors of subgenomic Hepatitis C Virus (HCV) RNA replication and pharmaceutical compositions of such compounds.

The novel compounds of this invention are novel purine and pyrimidine nucleoside derivatives listed as follows:

Compounds of formula I-a



wherein

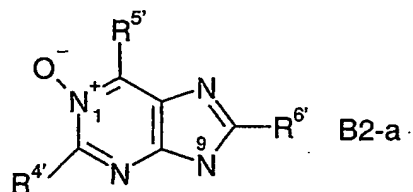
R^{1'} is hydroxy;

R^{2'} is hydroxy;

X' is O;

a', b', c', d' denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

B' signifies an oxidised purine base B2-a which is connected through the 9-nitrogen of formula



wherein

R^{4'} is hydrogen;

$R^{5''}$ is $NHR^{8''}$;

$R^{6''}$ is hydrogen;

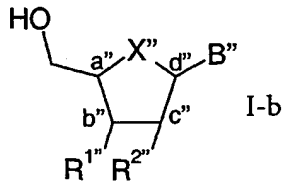
$R^{8''}$ is alkyl,

preferably wherein

- 5 $R^{8''}$ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, phenylmethyl (benzyl), 1-phenylethyl, 2-phenylethyl, 1(S)-methyl-2-phenylethyl, 1(R)-methyl-2-phenylethyl, 1-phenylpropyl, 2-phenylpropyl or 3-phenylpropyl;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

10 Compounds of formula I-b



wherein

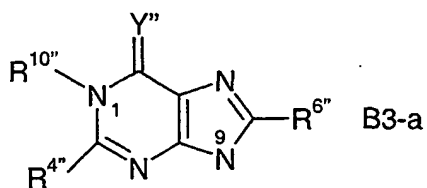
$R^{1''}$ is hydroxy;

$R^{2''}$ is hydroxy;

- 15 X'' is O;

a'', b'', c'', d'' denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

B'' signifies a purine base B3-a which is connected through the 9-nitrogen of formula



wherein

R⁴ is hydrogen;

R⁶ is hydrogen;

5 R¹⁰ is alkyl,

preferably wherein

R¹⁰ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl;

Y² is NR¹¹;

R¹¹ is alkyl,

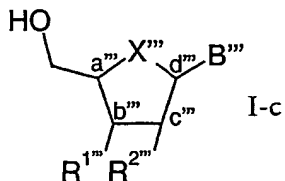
10 preferably wherein

R¹¹ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl, phenylmethyl (benzyl), 1-phenylethyl, 2-phenylethyl, 1(S)-methyl-2-phenylethyl, 1(R)-methyl-2-phenylethyl, 1-phenylpropyl, 2-phenylpropyl or 3-phenylpropyl;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

15

Compounds of formula I-c



wherein

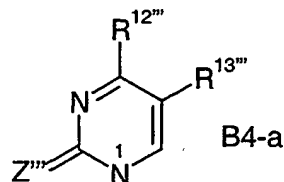
R^{1'''} is hydroxy;

R^{2'''} is hydroxy;

X''' is O;

a''', b''', c''', d''' denoting asymmetric carbon atoms and forming a D-
5 ribofuranosyl ring; and

group B''' signifies a pyrimidine base B4-a which is connected through the 1-nitrogen of formula



wherein

10 R^{12'''} is alkylthio or heterocyclyl,

preferably wherein

15 R^{12'''} is methylthio, ethylthio, n-propylthio, i-propylthio, n-butylthio, i-butylthio, tert.-butylthio or oxazolyl, isoxazolyl, furyl, tetrahydrofuryl, 2-thienyl, 3-thienyl, pyrazinyl, isothiazolyl, indolyl, didehydroindolyl, indazolyl, quinolinyl, pyrimidinyl, benzofuranyl, 1-pyrrolidinyl, 2-pyrrolidinyl, 3-pyrrolidinyl, 1-pyrrolyl, 2-pyrrolyl, triazolyl e.g. 1,2,3-triazolyl or 1,2,4-triazolyl, 1-pyrazolyl, 2-pyrazolyl, 4-pyrazolyl, benzotriazolyl, piperidinyl, morpholinyl (e.g. 4-morpholinyl), thiomorpholinyl (e.g. 4-thiomorpholinyl), thiazolyl, pyridinyl, dihydrothiazolyl, imidazolidinyl, pyrazolinyl, benzothienyl, piperazinyl, 1-imidazolyl, 2-imidazolyl, 4-imidazolyl, thiadiazolyl e.g. 1,2,3-thiadiazolyl, 1,2,3,4-tetrahydroquinoline, 1,2,3,4-tetrahydroisoquinoline, benzothiazolyl;

20 R^{13'''} is hydrogen, alkyl or halogen,

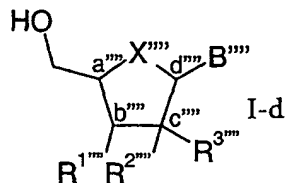
preferably wherein

25 R^{13'''} is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl or fluorine, chlorine, bromine or iodine;

Z''' is O;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

Compounds of formula I-d



5

wherein

R^{1'''} is hydrogen, halogen, hydroxy, alkyl, alkoxy, cyano or azido,

preferably wherein

R^{1'''} is hydrogen, fluorine, hydroxy, C₁₋₄-alkyl, C₁₋₄-alkoxy, cyano or azido,

10 more preferred wherein

R^{1'''} is hydrogen, fluorine, hydroxy, C₁₋₄-alkyl or C₁₋₄-alkoxy,

and most preferred wherein

R^{1'''} is hydroxy;

R^{2'''} and R^{3'''} represent fluorine;

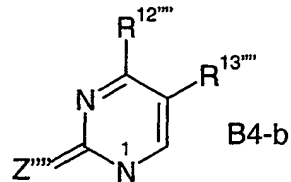
15 X''' is O or CH₂,

preferably wherein

X''' is CH₂;

a''', b''', c''', d''' denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

group B^{'''} signifies a pyrimidine base B4-b which is connected through the 1-nitrogen of formula



wherein

5 Z^{'''} is O;

R^{12'''} is NR^{7'''}R^{8'''},

preferably wherein

R^{12'''} is hydrogen, alkyl or halogen;

R^{13'''} is hydrogen, alkyl or halogen,

10 preferably wherein

R^{13'''} is hydrogen, C₁₋₄-alkyl or fluorine,

more preferred wherein

R^{13'''} is hydrogen, methyl, ethyl or fluorine,

and most preferred wherein

15 R^{13'''} is hydrogen;

R^{7'''} and R^{8'''} are independently of each other hydrogen or alkyl,

preferably wherein

R^{7'''} and R^{8'''} are independently of each other hydrogen or C₁₋₄-alkyl,

more preferred wherein

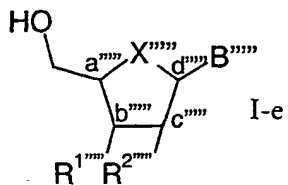
20 R^{7'''} and R^{8'''} are independently of each other hydrogen, methyl or ethyl,

and most preferred wherein

R^7 and R^8 are independently of each other hydrogen;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

5 Compounds of formula I-e



wherein

R^1 is alkoxy,

preferably wherein

10 R^1 is methoxy, ethoxy, n-propyloxy, i-propyloxy, n-butyloxy, i-butyloxy, tert.-butyloxy;

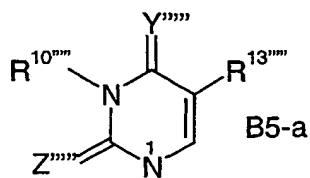
R^2 is hydrogen;

X is O;

a , b , c , d denoting asymmetric carbon atoms and forming a D-

15 ribofuranosyl ring; and

group B signifies a pyrimidine base B5-a which is connected through the 1-nitrogen of formula



wherein

$R^{10''''}$ is hydrogen;

$R^{13''''}$ is alkyl,

preferably wherein

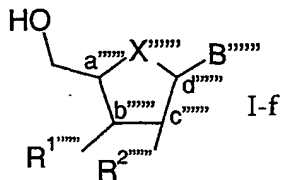
$R^{13''''}$ is methyl, ethyl, propyl, isopropyl, n-butyl, isobutyl, tert.-butyl;

5 Y'''' is O;

Z'''' is O;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

Compounds of formula I-f



10

wherein

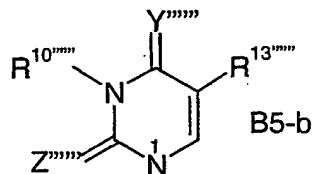
$R^{1''''}$ is hydroxy;

$R^{2''''}$ is hydroxy;

X'''' is O;

15 a'''' , b'''' , c'''' , d'''' denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

group B'''' signifies a pyrimidine base B5-b which is connected through the 1-nitrogen of formula



- 98 -

wherein

$R^{10\dots\dots\dots}$ is hydrogen;

$R^{13\dots\dots\dots}$ is halogen,

preferably wherein

5 $R^{13\dots\dots\dots}$ is fluorine, chlorine or bromine;

$Y^{\dots\dots\dots}$ is $NR^{11\dots\dots\dots}$;

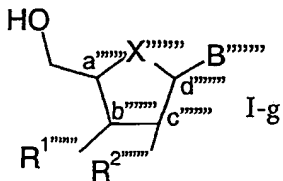
$R^{11\dots\dots\dots}$ is hydroxy;

$Z^{\dots\dots\dots}$ is O;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

10

Compounds of formula I-g



wherein

$R^{1\dots\dots\dots}$ is hydroxy;

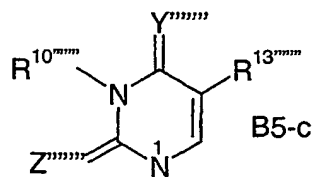
15 $R^{2\dots\dots\dots}$ is hydroxy;

$X^{\dots\dots\dots}$ is O;

$a^{\dots\dots\dots}$, $b^{\dots\dots\dots}$, $c^{\dots\dots\dots}$, $d^{\dots\dots\dots}$ denoting asymmetric carbon atoms and forming a L-ribofuranosyl ring; and

20 group B $^{\dots\dots\dots}$ signifies a pyrimidine base B5-c which is connected through the 1-nitrogen of formula

- 99 -



wherein

R¹⁰ is hydrogen;

R¹³ is hydrogen;

5 Y is NR¹¹;

R¹¹ is hydroxy;

Z is O;

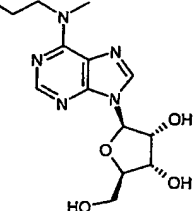
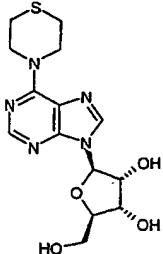
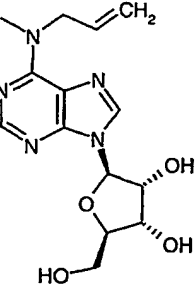
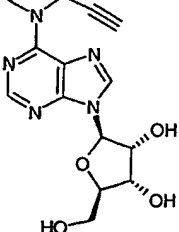
hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

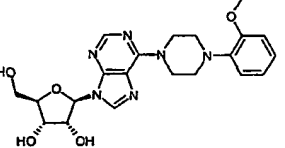
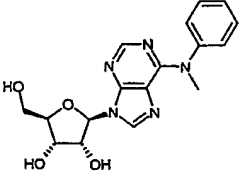
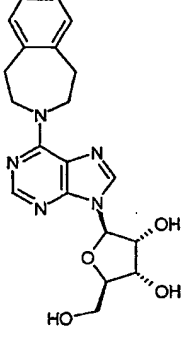
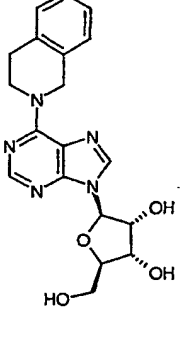
10

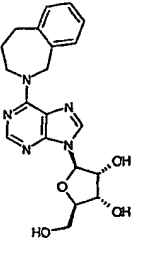
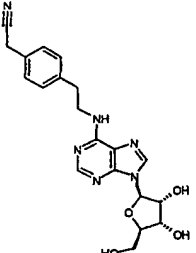
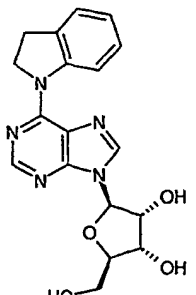
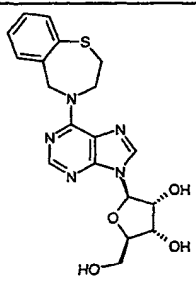
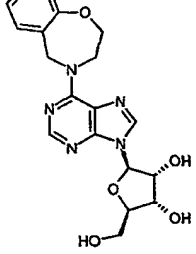
The terms as they are used for the novel purine and pyrimidine nucleoside derivatives are as defined above.

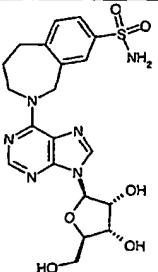
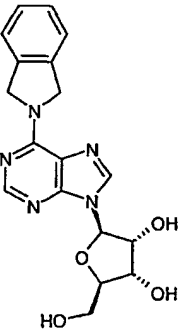
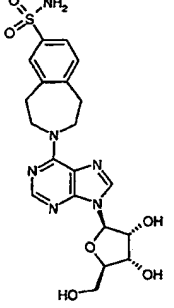
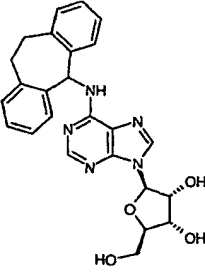
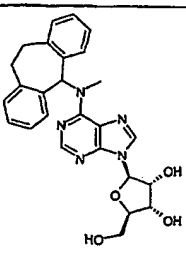
More preferred embodiments of compounds of formula I hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof, are listed in table 2:

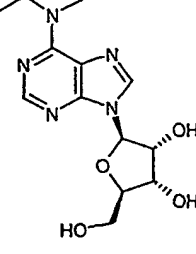
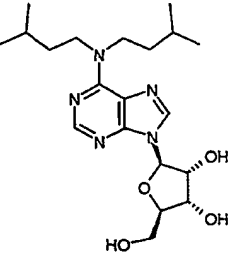
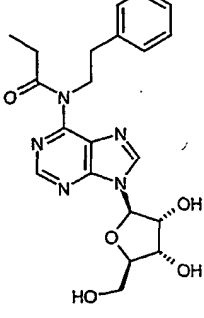
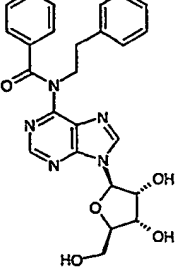
Table 2

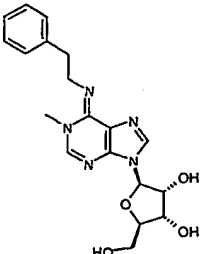
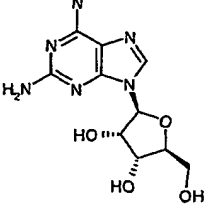
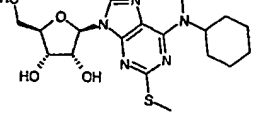
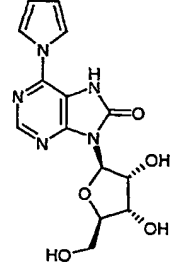
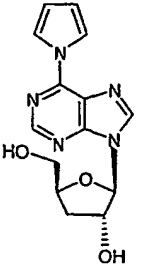
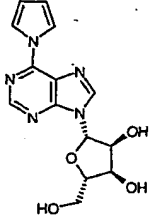
STRUCTURE	SYSTEMATIC NAME
	6-(N-Methylpropylamino)-9-(β -D-ribofuranosyl)purine
	9-(β -D-Ribofuranosyl)-6-(4-thiomorpholinyl)purine
	6-(N-Methyl-2-propenylamino)-9-(β -D-ribofuranosyl)purine
	6-(N-Methyl-2-propynylamino)-9-(β -D-ribofuranosyl)purine

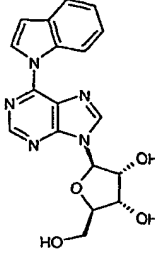
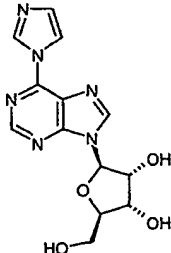
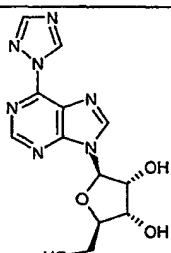
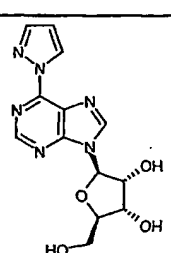
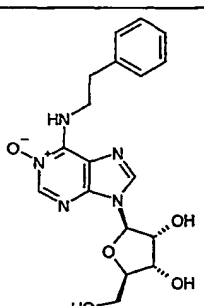
	<p>6-[4-(2-Methoxyphenyl)piperazinyl]-9-(β-D-ribofuranosyl)purine</p>
	<p>6-(N-Methylphenylamino)-9-(β-D-ribofuranosyl)purine</p>
	<p>9-(β-D-Ribofuranosyl)-6-(1,2,4,5-tetrahydro-3H-benzazepin-3-yl)purine</p>
	<p>9-(β-D-Ribofuranosyl)-6-(1,2,3,4-tetrahydro-2-isoquinolyl)purine</p>

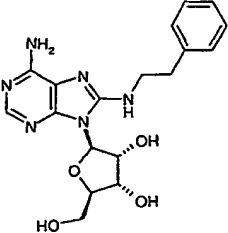
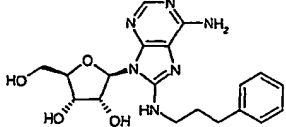
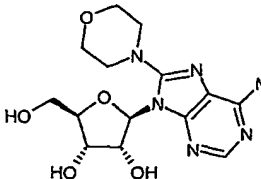
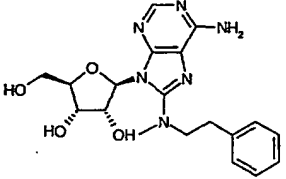
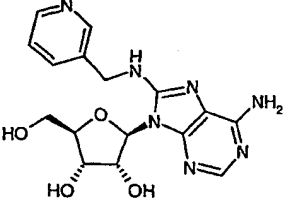
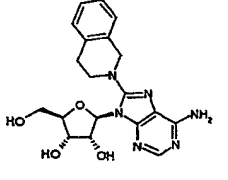
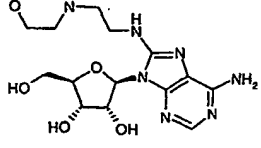
	9-(β -D-Ribofuranosyl)-6-(1,3,4,5-tetrahydro-2H-benzazepin-2-yl)purine
	6-[2-(4-Cyanomethylphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine
	6-(2,3-Dihydro-1-indolyl)- 9-(β -D-ribofuranosyl)purine
	9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl)purine
	9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl)purine

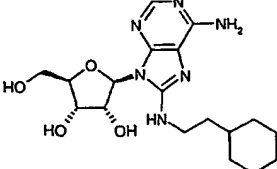
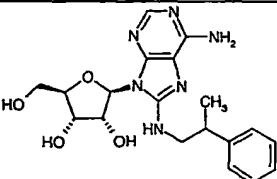
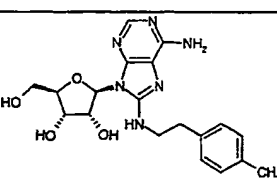
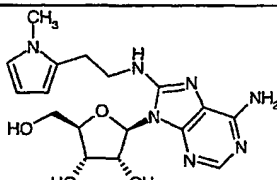
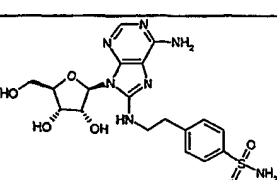
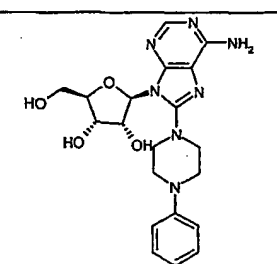
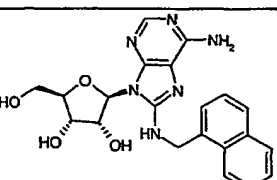
	<p>6-(8-Aminosulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl)-9-(β-D-ribofuranosyl)purine</p>
	<p>6-(2-Isoindolinyl)-9-(β-D-ribofuranosyl)purine</p>
	<p>6-(7-Aminosulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl)-9-(β-D-ribofuranosyl)purine</p>
	<p>6-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-ylamino)-9-(β-D-ribofuranosyl)purine</p>
	<p>6-[N-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-yl)methylamino]-9-(β-D-ribofuranosyl)purine</p>

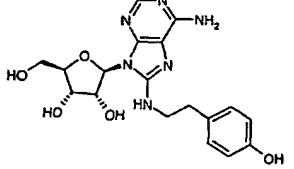
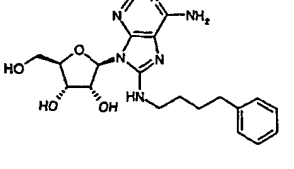
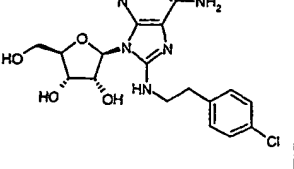
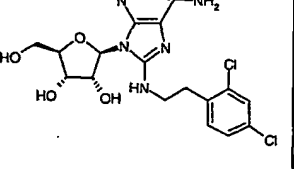
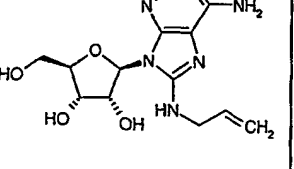
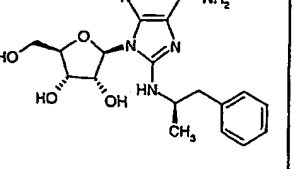
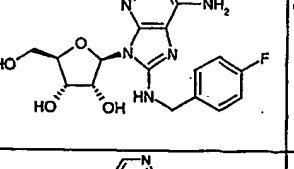
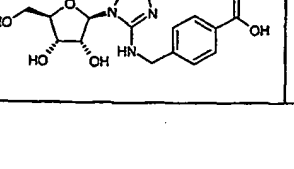
	<p>6-Ethylmethylamino- 9-(β-D-ribofuranosyl)purine</p>
	<p>6-bis-[(3-Methyl)butylamino]-9-(β-D-ribofuranosyl)purine</p>
	<p>6-[2-Phenyl-(N-propionyl)ethylamino]-9-(β-D-ribofuranosyl)purine</p>
	<p>6-(N-Benzoyl-2-phenylethylamino)-9-(β-D-ribofuranosyl)purine</p>

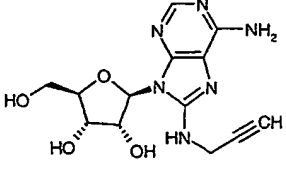
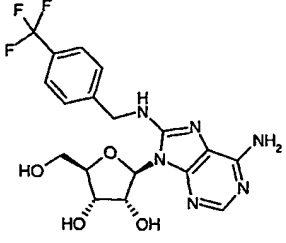
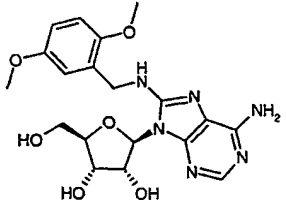
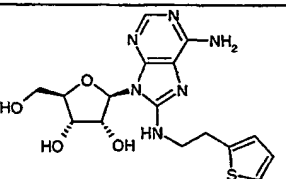
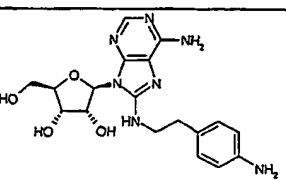
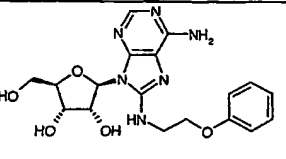
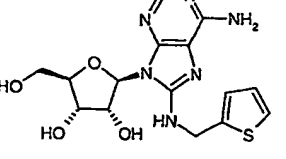
	1-Methyl-6-(2-phenylethylimino)-9-(β -D-ribofuranosyl)purine
	2-Amino-6-methylamino-9-(β -L-ribofuranosyl)purine
	6-[(N-Cyclohexyl)methylamino]-2-methylthio-9-(β -D-ribofuranosyl)purine
	6-(1-Pyrrolyl)-9-(β -D-ribofuranosyl)purin-8-(7H)-one
	9-(3-Deoxy- β -D-ribofuranosyl)-6-(1-pyrrolyl)purine
	6-(1-Pyrrolyl)-9-(β -L-ribofuranosyl)purine

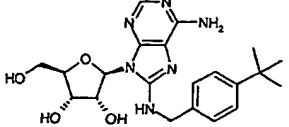
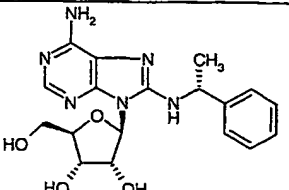
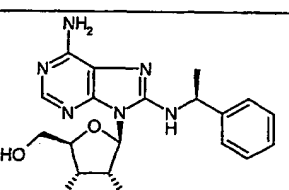
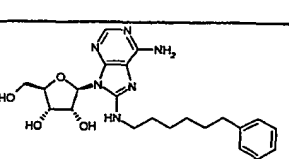
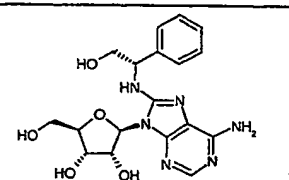
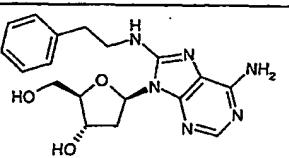
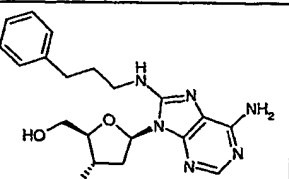
	6-(1-Indolyl)-9-(β -D-ribofuranosyl)purine
	6-(1-Imidazolyl)-9-(β -D-ribofuranosyl)purine
	9-(β -D-Ribofuranosyl)-6-(1,2,4-triazol-1-yl)purine
	6-(1-Pyrazolyl)- 9-(β -D-ribofuranosyl)purine
	6-(2-Phenylethylamino)- 9-(β -D-ribofuranosyl)purine-1-oxide

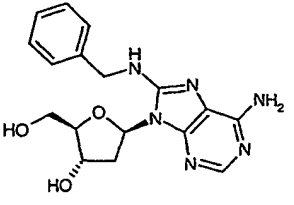
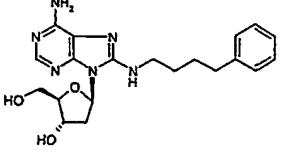
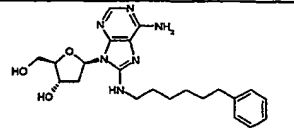
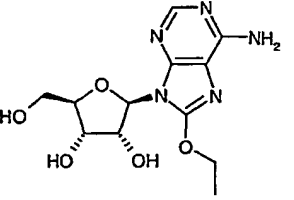
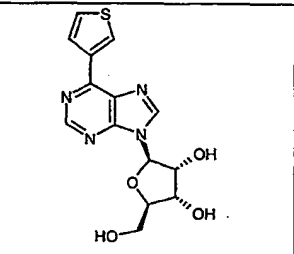
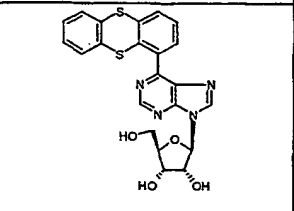
	8-(2-Phenylethylamino)adenosine
	8-(3-Phenylpropylamino)adenosine
	8-(4-Morpholinyl)adenosine
	8-(N-Methyl-2-phenylethylamino)adenosine
	8-(3-Pyridylmethylamino)adenosine
	8-(1,2,3,4-Tetrahydro-2-isoquinolyl)adenosine
	8-[2-(4-Morpholinyl)ethylamino]adenosine

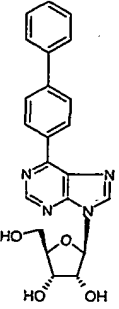
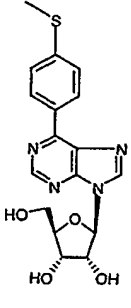
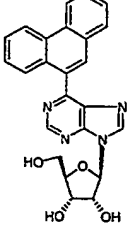
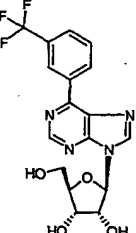
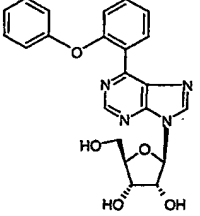
	8-(2-Cyclohexylethylamino)adenosine
	8-(2(R,S)-Phenylpropylamino)adenosine
	8-[2-(4-Methylphenyl) ethylamino]adenosine
	8-[2-(1-Methyl-2-pyrrolyl) ethylamino]adenosine
	8-[2-(4-Aminosulphonylphenyl) ethylamino]adenosine
	8-(4-Phenyl-1-piperazinyl)adenosine
	8-(1-Naphthylmethylamino)adenosine

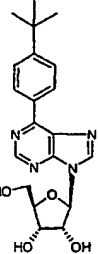
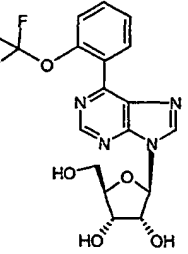
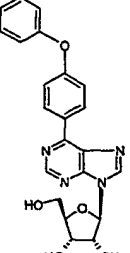
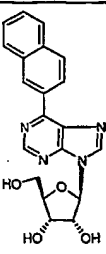
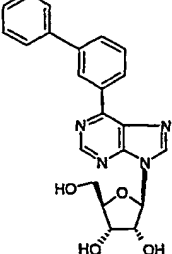
	8-[2-(4-Hydroxyphenyl)ethylamino]adenosine
	8-(4-Phenylbutylamino)adenosine
	8-[2-(4-Chlorophenyl)ethylamino]adenosine
	8-[2-(2,4-Dichlorophenyl)ethylamino]adenosine
	8-(2-Propenylamino)adenosine
	8-(1(R)-Methyl-2-phenylethylamino)adenosine
	8-(4-Fluorobenzylamino)adenosine
	8-[(4-Hydroxycarbonyl)benzylamino]adenosine

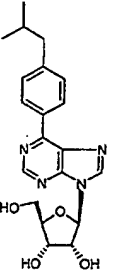
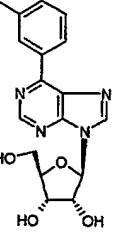
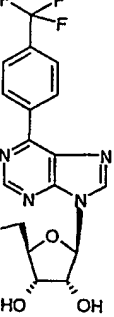
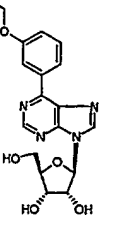
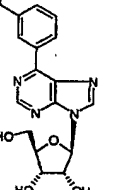
	8-(2-Propynylamino)adenosine
	8-[(4-Trifluoromethyl)benzylamino]adenosine
	8-[(2,5-Dimethoxy)benzylamino]adenosine
	8-[2-(2-Thienyl)ethylamino]adenosine
	8-[2-(4-Aminophenyl)ethylamino]adenosine
	8-(2-Phenoxyethylamino)adenosine
	8-[(2-Thienyl)methylamino]adenosine

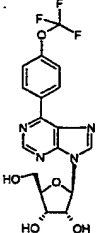
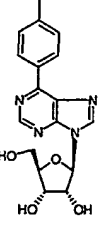
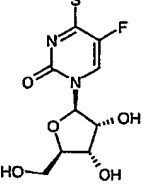
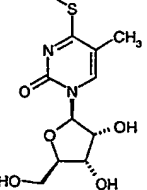
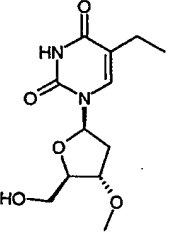
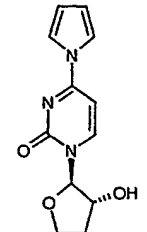
	8-[(4-tert-Butyl)benzylamino]adenosine
	8-(1(R)-Phenylethylamino)adenosine
	8-(1(S)-Phenylethylamino)adenosine
	8-(6-Phenylhexylamino)adenosine
	8-[2-Hydroxy-1(S)-phenylethylamino]adenosine
	2'-Deoxy-8-(2-phenylethylamino)adenosine
	2'-Deoxy-8-(3-phenylpropylamino)adenosine

	8-Benzylamino-2'-deoxyadenosine
	2'-Deoxy-8-(4-phenylbutylamino)adenosine
	2'-Deoxy-8-(6-phenylhexylamino)adenosine
	8-Ethoxyadenosine
	9-(β -D-Ribofuranosyl)-6-(3-thienyl)purine
	9-(β -D-Ribofuranosyl)-6-(1-thianthrenyl)purine

	6-(4-Biphenylyl)-9-(β -D-ribofuranosyl)purine
	6-(4-Methylthiophenyl)-9-(β -D-ribofuranosyl) purine
	6-(9-Phenanthrenyl)-9-(β -D-ribofuranosyl)purine
	9-(β -D-Ribofuranosyl)-6-(3- trifluoromethylphenyl)purine
	6-(2-Phenoxyphenyl)-9-(β -D-ribofuranosyl)purine

	6-(4-tert-Butylphenyl)-9-(β -D-ribofuranosyl)purine
	9-(β -D-Ribofuranosyl)-6-(2-trifluoromethoxyphenyl)purine
	6-(4-Phenoxyphenyl)-9-(β -D-ribofuranosyl)purine
	6-(2-Naphthyl)-9-(β -D-ribofuranosyl)purine
	6-(3-Biphenylyl)-9-(β -D-ribofuranosyl)purine

	6-[4-(2-Methylpropyl)phenyl]-9-(β -D-ribofuranosyl)purine
	6-(3-Fluorophenyl)-9-(β -D-ribofuranosyl)purine
	9-(β -D-Ribofuranosyl)-6-(4-trifluoromethylphenyl)purine
	6-(3-Ethoxyphenyl)-9-(β -D-ribofuranosyl)purine
	6-[3-(1-Methyl)ethylphenyl]-9-(β -D-ribofuranosyl)purine

	9-(β -D-Ribofuranosyl)-6-(4-trifluoromethoxyphenyl)purine
	6-(4-Ethylphenyl)-9-(β -D-ribofuranosyl)purine
	5-Fluoro-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
	5-Methyl-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
	2',3'-Dideoxy-5-ethyl-3'-methoxyuridine
	4-(1-Pyrrrolyl)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one

	4-Oximino-1-(β -L-ribofuranosyl)pyrimidin-2(1H)-one
	5-Fluoro-4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one
	4-Amino-1-(2,2-difluoro-3-hydroxy-4-hydroxymethyl-cyclopentyl)-1H-pyrimidin-2-one

5 The novel purine and pyrimidine nucleoside derivatives of formula I have been shown to be inhibitors of subgenomic Hepatitis C Virus replication in a hepatoma cell line. These compounds have the potential to be efficacious as antiviral drugs for the treatment of HCV infections in human. Accordingly, the present novel purine and pyrimidine nucleoside derivatives of formula I are therapeutically active substances in the treatment of HCV infections in human and can be used as medicaments for the treatment of such disease.

10 The novel purine and pyrimidine nucleoside derivatives of formula I can as well be used as medicaments, especially for treating immune mediated conditions or diseases, viral diseases, bacterial diseases, parasitic diseases, inflammatory diseases, hyperproliferative vascular diseases, tumors, and cancer.

15 In particular, compounds of the present invention and pharmaceutical compositions containing the same are useful as chemotherapeutic agents, inhibitors of viral replication and modulators of the immune system, and can be used for the treatment of viral diseases such as retroviral infections and hepatitis C virus infections (either alone or in combination with other antiviral agents such as interferon or derivatives thereof, such as conjugates with polyethylene glycol).

20 They can be used alone, or in combination with other therapeutically active agents, for example, an immunosuppressant, a chemotherapeutic agent, an anti-viral agent, an antibiotic, an anti-parasitic agent, an anti-inflammatory agent, an anti-fungal agent and/or an anti-vascular hyperproliferation agent.

25 Any functional (i.e. reactive) group present in a side-chain may be protected, with the protecting group being a group which is known per se, for example, as described in "Protective Groups in Organic Synthesis", 2nd Ed., T.W. Greene and P.G.M. Wuts, John Wiley & Sons, New York, NY, 1991. For example, an amino group can be protected by tert.-butyloxycarbonyl (BOC) or benzyloxycarbonyl (Z).

30 The compounds of this invention may contain one or more asymmetric carbon atoms and may therefore occur as racemates and racemic mixtures, single enantiomers, diastereomeric mixtures and individual diastereomers. Furthermore, where a compound of the invention contains an olefinic double bond, this can have the (E) or (Z) configuration. Also, each chiral center may be of the R or S

configuration. All such isomeric forms of these compounds are embraced by the present invention.

Compounds of formula I which are acidic can form pharmaceutically acceptable salts with bases such as alkali metal hydroxides, e.g. sodium hydroxide and potassium hydroxide; alkaline earth metal hydroxides, e.g. calcium hydroxide, barium hydroxide and magnesium hydroxide, and the like; with organic bases e.g. N-ethyl piperidine, dibenzylamine, and the like. Those compounds of formula I which are basic can form pharmaceutically acceptable salts with inorganic acids, e.g. with hydrohalic acids such as hydrochloric acid and hydrobromic acid, sulphuric acid, nitric acid and phosphoric acid, and the like, and with organic acids, e.g. with acetic acid, tartaric acid, succinic acid, fumaric acid, maleic acid, malic acid, salicylic acid, citric acid, methanesulphonic acid and p-toluene sulphonic acid, and the like. The formation and isolation of such salts can be carried out according to methods known in the art.

Also part of the present invention are known purine and pyrimidine nucleoside derivatives for use in medicine, especially for use in the treatment of an Hepatitis C Virus (HCV) infection, where no medical use for those compounds is previously known, and pharmaceutical compositions containing the same.

Assay Method: The activity of the compounds was assayed using an adaptation of the method reported by Lohmann et al [V. Lohmann et al., *Science*, 1999, 285, 110-113].

HCV Replicon Assay:

5 The HCV replicon-containing cell line is used for the identification of small molecules that are able to inhibit the replication of the replicon RNA. Since the replicon RNA replication mimics the replication of the HCV RNA in infected hepatocytes, it is believed that those small molecules that have the above property are interesting for further development as anti-HCV drugs.

10 The inhibition of the HCV replicon RNA replication will lead to a decrease of the replicon RNA in the cell, which can be measured using a method that specifically quantifies this RNA.

Northern blot: One method for quantification of this RNA uses the standard Northern blot known to any person skilled in the art.

15 Kinetic PCR: A second assay for the quantification of replicon RNA is based on the amplification of the replicon RNA that remains in the cell, after incubation of the cells with a proper concentration of the small molecules. This method involves the reverse transcription of the replicon RNA to the corresponding complementary DNA (cDNA), followed by amplification of the cDNA using the
20 Taqman Kinetic PCR technology (PE Biosystems). This consists of hybridisation of the cDNA with a complementary reporter oligonucleotide (probe), containing a combined fluorescent dye and a quencher dye. Amplification of the DNA sequence containing the hybridised reporter probe, using flanking oligonucleotide primers will lead to the separation of the fluorescent dye from the quencher dye. This will
25 result in an increase of the fluorescence during each amplification cycle.

The neomycin phosphotransferase gene sequence that is present in the replicon RNA was chosen for amplification using specifically designed oligonucleotide primers. To control for (a) cell number that can vary depending on the toxicity or cytostatic effect of the small molecules, and (b) for errors during
30 total RNA extraction, amplification of the host β -actin gene is used for normalisation.

The accumulation of the PCR products during the reaction is monitored directly by measuring the increase in fluorescence of the reporter dye. The amount

of HCV replicon RNA (and β -actin RNA) originally present in the total RNA extracted from the cells is then expressed as a threshold cycle, e.g. the cycle at which there is a statistically significant increase in the fluorescence above the background.

5 For this procedure, HCV replicon-containing human hepatoma Huh7 cells
 (9-13) in growth medium (DMEM) containing 5% FCS are plated in a 96-well
 plate at 5×10^3 cells per well, and the plate incubated overnight. 24 hours later,
 different dilutions in 0.1ml growth medium of chemical compounds were added to
 the wells, and the plate further incubated at 37°C for three days. Total RNA coming
10 from each well is extracted using the RNeasy™ procedure (Qiagen manufacturer
 instructions), and the total RNA is eluted in a final volume of 0.13ml. Next, a 2 μ l
 sample of the total RNA is used for conversion into cDNA using a reverse
 transcription (RT) step. A RT mastermix containing 1 μ l 10x Taqman RT buffer,
 2.2 μ l 25mM MgCl₂ (5.5mM final conc.), 2 μ l dNTP mix (500 μ M each), 0.5 μ l
15 random hexamer primers (2.5 μ M), 0.2 μ l RNase inhibitor (0.4u/ μ l), 0.25 μ l RT
 (1.25u/ μ l), 1.85 μ l H₂O, was distributed in a 96-well plate and 2 μ l total RNA was
 added to each well. The RT reaction is performed by incubation of the plate 10 min
 at 25°C, 30 min at 48°C, 5 min at 95°C and cooling to 4°C. The cDNA samples are
 then stored at -20°C or directly used for the PCR reaction. For the PCR reaction,
20 the cDNA is diluted by addition of 90 μ l water, and 10 μ l of each diluted cDNA
 sample is added in duplicate to each well of a 96-well optical plate containing
 12.5 μ l Taqman Universal PCR mix (PE Biosystems), 1.25 μ l 20x Replicon
 probe/primer mix (Primers 300nM, Probe 100nM), 1.25 μ l 20x β -actin
 probe/primer mix (PDAR PE Biosystems). A standard curve is generated for each
25 plate by including in duplicate five 3-fold dilutions of cDNA derived from total
 RNA extracted from 9-13 cell that were incubated in the absence of chemical
 compounds. A negative control is included in the plate by omitting the cDNA
 sample (no template control). Each well of the optical plate is secured with a lid
 and the plate is mixed. The plate is centrifuged for a few seconds at 3000 rpm to
 ensure contents are at the bottom of each well. The plate is then inserted into the
 7700 Kinetic PCR machine and the reaction started using the default settings.
30

35 The concentration of the drug (IC_{50}) required to reduce replicon RNA levels
 by 50% relative to the untreated 9-13 cell control value, can be calculated from the
 plot of percentage replicon RNA reduction vs. drug concentration.

Renilla Luciferase reporter: A third assay is based on the idea of using a reporter as a simple readout for intracellular HCV replicon RNA level. For this purpose the Renilla luciferase gene was introduced into the first open reading frame of a replicon construct NK5.1 (Krieger *et al.*, J. Virol. 75:4614), immediately after the internal ribosome entry site (IRES) sequence, and fused with the neomycin phosphotransferase (NPTII) gene via a self-cleavage peptide 2A from foot and mouth disease virus (Ryan & Drew, EMBO Vol 13:928-933). After *in vitro* transcription the RNA was electroporated into human hepatoma Huh7 cells, and G418-resistant colonies were isolated and expanded. Stably selected cell line 2209-23 was shown to contain replicative HCV subgenomic RNA, and the activity of Renilla luciferase expressed by the replicon reflects its RNA level in the cells.

For the assay procedure, Renilla Luciferase HCV replicon cells (2209-23) that cultured in Dulbecco's MEM (GibcoBRL cat no. 31966-021) with 5% fetal calf serum (FCS) (GibcoBRL cat no. 10106-169) were plated onto a 96-well plate at 5000 cells per well, and incubated overnight. Twenty-four hours later, different dilutions of chemical compounds in the growth medium were added to the cells, which were then further incubated at 37°C for three days. The assay was carried out in duplicate plates, one in opaque white and one in transparent, in order to measure the activity and cytotoxicity of a chemical compound in parallel ensuring the activity seen is not due to reduction on cell proliferation.

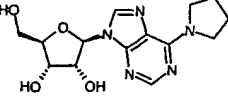
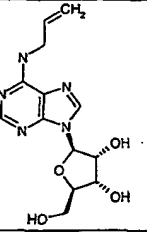
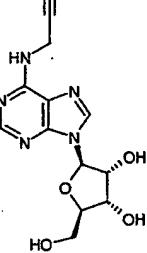
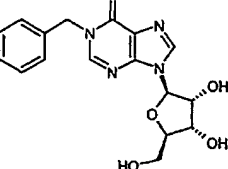
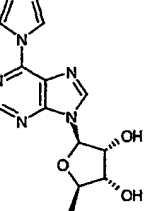
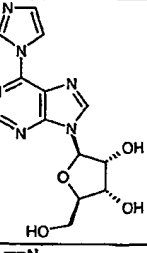
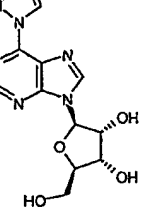
At the end of the incubation time, the cells in the white plate were harvested and luciferase activity was measured by using a Dual-Luciferase reporter assay system (Promega cat no. E1960). All the reagents described in the following paragraph were included in the manufacturer's kit, and the manufacturer's instructions were followed for preparations of the reagents. Briefly, the cells were washed twice with 200µl PBS (phosphate buffered saline; pH 7.0) per well and lysed with 25µl of 1x passive lysis buffer prior to incubation at room temperature for 20 min. One hundred microlitre of LAR II reagent was added to each well. The plate was then inserted into the LB 96V microplate luminometer (MicroLumatPlus, Berthold), and 100 µl of Stop & Glo reagent was injected into each well by the machine and the signal measured using a 2-second delay, 10-second measurement programme. The IC₅₀, the concentration of the drug required for reducing the replicon level by 50% in relation to the untreated cell control value, can be calculated from the plot of the percentage reduction of the luciferase activity vs. drug concentration.

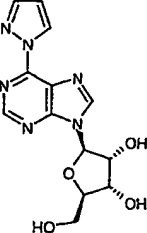
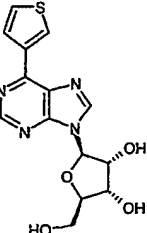
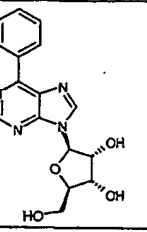
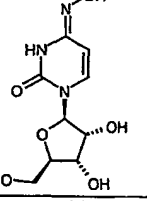
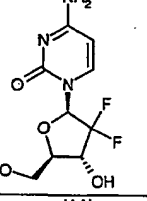
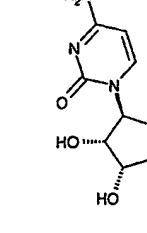
Biological Test results:

Compounds were tested for inhibition of HCV replicon RNA replication using the above assay. Examples of the results are shown in the following table:

5

Example	Structure	Name	IC ₅₀ (μM)
1		6-Dimethylamino-9-(β-D-ribofuranosyl)purine	0.6
7		Adenosine-1-oxide	2
10		8-Bromoadenosine	3.6
16		6-Methylthio-9-(β-D-ribofuranosyl)purine	0.08
19		6-Chloro-9-(β-D-ribofuranosyl)purine	14
24		5-Fluorouridine	1.4
57		9-(β-D-Ribofuranosyl)-6-(1,2,3,4-tetrahydro-2-isoquinolyl)purine	0.1

77		6-(1-Pyrrolidinyl)-9-(β-D-ribofuranosyl)purine	2.6
80		6-(2-Propenyl)amino-9-(β-D-ribofuranosyl)purine	5.7
81		6-(2-Propynyl)amino-9-(β-D-ribofuranosyl)purine	3.8
93		1-Benzyl-6-imino-9-(β-D-ribofuranosyl)purine	4.5
105		6-(1-Pyrrolyl)-9-(β-D-ribofuranosyl)purine	0.1
111		6-(1-Imidazolyl)-9-(β-D-ribofuranosyl)purine	6.2
112		9-(β-D-Ribofuranosyl)-6-(1,2,4-triazol-1-yl)purine	4.4

113		6-(1-Pyrazolyl)-9-(β-D-ribofuranosyl)purine	4.4
181		9-(β-D-Ribofuranosyl)-6-(3-thienyl)purine	0.05
182		6-Phenyl-9-(β-D-ribofuranosyl) purine	0.1
239		4-Oximino-1-(β-D-ribofuranosyl)pyrimidin-2(1H)-one	1.3
243		1-(2-Deoxy-2,2-difluoro-β-D-erythropentofuranosyl)cytosine	0.07
244		L-Cytidine	10

- 126 -

245		4-Amino-1-(2,2-difluoro-3-hydroxy-4-hydroxymethylcyclopentyl)-1H-pyrimidin-2-one	2
246		4-Amino-1(R)-(2(S),3(R))-dihydroxy-4(R)-hydroxymethylcyclopentyl)-1H-pyrimidin-2-one	0.4
247		1-(β-D-Xylofuranosyl)cytosine	3.7
249		1-(3-Deoxy-3-fluoro-β-D-xylofuranosyl)cytosine	10.4
252		6-Ethylamino-9-(β-D-ribofuranosyl)purine	14
253		6-Propylamino-9-(β-D-ribofuranosyl)purine	7

Compounds 246, 247, 249, 252 and 253 were tested in the Renilla luciferase assay.

Dosing for the human body with compounds of formula I:

5

The compounds according to the invention may be employed alone or in combination with other therapeutic agents for the treatment of hepatitis C virus infections.

5 The compound of formula I whether administered alone or in combination with other therapeutic agents may be administered orally in capsule, tablet or liquid form. Other types of administration could also be contemplated such as nasal spray, transdermally, by suppository, by sustained release dosage form and by pulmonary inhalation, as long as adequate dosages are delivered without destroying the active ingredient.

10 The amount of the compound of formula I required for the treatment of hepatitis C virus infections will depend on a number of factors including the severity of the disease and the identity, sex and weight of the recipient and will ultimately be at the discretion of the attendant physician. In general, however, a suitable effective dose is in the range of 0.05 to 100mg per kilogram of body weight of the recipient per day, preferably in the range 0.1 to 50mg per kilogram of body weight per day and most preferably in the range of 0.5 to 20mg of body weight per day. An optimum dose is about 2 to 16mg per kilogram body weight per day. The desired dose is preferably presented as two, three, four, five , six or more sub-doses administered at appropriate intervals throughout the day. These sub-doses may be administered in unit dosage forms, for example, containing from 1 to 1500mg, preferably from 5 to 1000mg, most preferably from 10 to 700mg of active ingredient per unit dosage form.

20 Combination therapies comprise the administration at least one compound of formula I or a physiologically functional derivative and at least one other physiologically acceptable agent. The active ingredient(s) and physiologically acceptable agent(s) may be administered together or separately and when administered separately this may occur simultaneously or sequentially in any order. The amounts of the active ingredient(s) and physiologically acceptable agent(s) and the relative timings of administration will be selected in order to achieve the desired combined therapeutic effect. Preferably the combination therapy involves the administration of one compound of formula I or a physiologically functional derivative and interferon alpha. The interferon alpha administered is preferably selected from interferon alpha 2a, interferon alpha 2b, a consensus interferon, a purified interferon alpha product or a pegylated interferon alpha 2a or a pegylated interferon alpha 2b. Preferably the amount of interferon alpha administered is from 2 to 10 million IU per week on a weekly, TIW, QOD or daily basis. The preferred method of administering the interferon alpha or pegylated interferon alpha formulations is parenterally, preferably by subcutaneous, IV, or IM injection.

It is preferable to administer the compound of formula I as a pharmaceutical formulation. The formulations of the present invention comprise at least one active ingredient of formula I together with one or more pharmaceutically acceptable excipients and optionally one or more other therapeutic agents.

- 5 Formulations for oral administration may be capsules, cachets or tablets each containing a predetermined amount of active ingredient(s) may be prepared by any method well known in the art of pharmacy. As well as the active ingredients(s) the oral formulation may contain a binder (for example povidone, gelatin, hydroxypropylmethyl cellulose), a lubricant, inert diluent, preservative,
- 10 disintegrant (for example sodium starch glycollate, cross-linked povidone, cross-linked sodium carboxymethyl cellulose) or a dispersing agent. Formulations for oral use may also include buffering agents to neutralise stomach acidity.

Example:

- Tablets containing the following ingredients may be produced in a
15 conventional manner:

Ingredient	per tablet
Compound of formula I	100mg
Lactose	131mg
Microcrystalline cellulose	60mg
20 Croscarmellose sodium	6mg
Magnesium stearate	<u>3mg</u>
Tablet weight	300mg

The following examples for the preparation of compounds of formula I illustrate the present invention. The known compounds of formula I are mostly commercially available (the supplier is indicated) or can be synthesised according the below procedure:

5

Example 1: 6-Dimethylamino-9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No. D2754.

Example 2: 6-(1(S)-Methyl-2-phenylethylamino)-9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No. P7665.

10 Example 3: 3'-Deoxyadenosine, Sigma-Aldrich Company Ltd., Cat. No. C3394.

Example 4: 6-(2-Phenylethylamino)- 9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd. Cat. No. P2673.

Example 5: 6-Cyclohexylamino-9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No.C9901.

15 Example 6: 2-Chloroadenosine, Aldrich Chemical Company, Cat. No. 86,186-3.

Example 7: Adenosine-1-oxide, Sigma-Aldrich Company Ltd., Cat. No.A8540.

Example 8: 9-(β -D-Ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No. P9278.

Example 9: 3'-Deoxyguanosine, Sigma-Aldrich Company Ltd., Cat. No. D7285.

20 Example 10: 8-Bromoadenosine, Aldrich Company Ltd., Cat. No.12,750-7.

Example 11: 8-Bromo-2'-deoxyadenosine, Maybridge Chemical Company, Cat. No.BTB14107.

Example 12: 8-Bromoguanosine, Sigma-Aldrich Company Ltd., Cat. No. B1893.

Example 13: 6-Thioguanosine, Sigma-Aldrich Company Ltd., Cat. No. M6625.

25 Example 14: Inosine, Sigma-Aldrich Company Ltd., Cat. No. I1024.

Example 15: 6-Thioinosine, Sigma-Aldrich Company Ltd., Cat. No. M7250.

- 130 -

- Example 16: 6-Methylthio-9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No. M4002.
- Example 17: L-Inosine, Penta, Cat. No. 09-02700.
- Example 18: 8-Bromoinosine, Sigma-Aldrich Company Ltd., Cat. No. B4004.
- 5 Example 19: 6-Chloro-9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No. C8276.
- Example 20: 2-Amino-6-chloro-9-(β -D-ribofuranosyl)purine, Sigma-Aldrich Company Ltd., Cat. No. A4634.
- 10 Example 21: 2'-Deoxy-5-fluorouridine, Sigma-Aldrich Company Ltd., Cat. No. F0503.
- Example 22: 1-(β -D-Arabinofuranosyl)-5-fluorouracil, George-Uhe Company Inc., Cat. No. 000265.
- Example 23: 4-Thiouridine, Sigma-Aldrich Company Ltd., Cat. No. T4509.
- Example 24: 5-Fluorouridine, Sigma-Aldrich Company Ltd., Cat. No. F5130.
- 15 Example 25: 5-Bromouridine, Sigma-Aldrich Company Ltd., Cat. No. B9752.
- Example 26: 3-Methyluridine, Sigma-Aldrich Company Ltd., Cat. No. M4129.
- Example 27: 5-Methyluridine, Sigma-Aldrich Company Ltd., Cat. No. M8905.
- Example 28: 1-(β -D-Arabinofuranosyl)uracil, Sigma-Aldrich Company Ltd., Cat. No. M8905.
- 20 Example 29: 1-(β -D-Arabinofuranosyl)-5-methyluracil, Sigma-Aldrich Company Ltd., Cat. No. T3766.
- Example 30: 1-(β -D-Arabinofuranosyl)-5-iodouracil, George-Uhe Company Inc., Cat. No. 000322.
- Example 31: 3'-Deoxy-5-methyluridine, Berry, Cat. No. PY7260.
- 25 Example 32: 5-Fluorocytidine, ICN Biomedicals Inc., Cat. No. 151156.

- 131 -

Example 33: 1-(β -D-Arabinofuranosyl)-5-fluorocytosine, Sigma-Aldrich Company Ltd., Cat. No. F3504.

Example 34: 5-Methylcytidine, Sigma-Aldrich Company Ltd., Cat. No. M4524.

Example 35: 2',3'-Dideoxycytidine, Sigma-Aldrich Company Ltd., Cat. No. D5782.

5 Example 36: N4-Acetylcytidine, Sigma-Aldrich Company Ltd., Cat. No. A7766.

Example 37: 3'-Deoxycytidine, Sigma-Aldrich Company Ltd., Cat. No. D5179.

Example 38

10 0.25g of 6-chloro-9-(β -D-ribofuranosyl)purine and 0.7g of N-methylpropylamine in 5ml of anhydrous ethanol were heated at reflux temperature for 1 hour. After cooling to room temperature, the solution was concentrated under reduced pressure and the mixture purified by flash column chromatography on silica gel using methanol/dichloromethane (10:90) as the eluent, to give 0.04g of 15 6-(N-methylpropylamino)-9-(β -D-ribofuranosyl)purine as a light yellow solid; mass spectrum (ESI) 324 [M+H]⁺.

Example 39

20 Reaction of 6-chloro-9-(β -D-ribofuranosyl)purine with thiomorpholine in an analogous manner to that described in example 38, gave 9-(β -D-ribofuranosyl)-6-(4-thiomorpholinyl)purine as a light brown solid; mass spectrum (ESI) 354 25 [M+H]⁺.

Example 40

Reaction of 6-chloro-9-(β -D-ribofuranosyl)purine with N-methylallylamine in an analogous manner to that described in example 38, gave 6-(N-methyl-2-propenylamino)-9-(β -D-ribofuranosyl)purine as an off-white solid; mass spectrum (ESI) 322 [M+H]⁺.

Example 41

Reaction of 6-chloro-9-(β -D-ribofuranosyl)purine and N-methylpropargylamine in an analogous manner to that described in example 38, gave 6-(N-methyl-2-propynylamino)-9-(β -D-ribofuranosyl)purine as an off-white solid; mass spectrum (ESI) 320 [M+H]⁺.

Also in a manner analogous to that described in example 38 starting with 6-chloro-9-(β -D-ribofuranosyl)purine and the appropriate amine were prepared the following examples:

10

Example 42: 6-(4-Morpholinyl)-9-(β -D-ribofuranosyl)purine, (K. Kikugawa et al, J. Med. Chem., 1972, 15, 387).

Example 43: 6-Diethylamino-9-(β -D-ribofuranosyl)purine, (Walsh et al, J.Amer.Chem.Soc., 1967, 89, 6221).

15

Example 44: 6-(1(R,S)-Phenylethylamino)-9-(β -D-ribofuranosyl)purine, (S. Kusachi et al, J. Med. Chem., 1985, 28, 1636).

Example 45: 6-(1-Benzyl-1-methylethylamino)-9-(β -D-ribofuranosyl)purine, (S. Kusachi et al, J. Med. Chem., 1985, 28, 1636).

20

Example 46: 6-(3-Phenylpropylamino)-9-(β -D-ribofuranosyl)purine, (S. Kusachi et al, J. Med. Chem., 1985, 28, 1636).

Example 47: 9-(β -D-Ribofuranosyl)-6-[2-(2-thienyl)ethylamino]purine, (S. Kusachi et al, J. Med. Chem., 1985, 28, 1636).

Example 48: 6-Dibenzylamino-9-(β -D-ribofuranosyl)purine, (Endo and Zemlicka, J. Org. Chem., 1979, 44, 3652).

25

Example 49: 6-Hexylamino-9-(β -D-ribofuranosyl)purine, (S. Kusachi et al, J. Med. Chem., 1985, 28, 1636).

Example 50: 6-(3-Pyridylmethylamino)-9-(β -D-ribofuranosyl)purine, (Kissmann and Weiss, J. Org. Chem., 1956, 21, 1053).

Example 51: 6-[4-(4-Fluorophenyl)-1,2,5,6-tetrahydropyridyl]-9-(β -D-ribofuranosyl)purine.

Example 52: 6-[4-(2-Methoxyphenyl)piperazinyl]-9-(β -D-ribofuranosyl)purine.

Example 53: 6-[2-(3-Indolyl)ethylamino]-9-(β -D-ribofuranosyl)purine, (Shikita et al, Chem. Pharm.Bull., 1974, 22, 1410).

Example 54: 6-[2-(4-Chlorophenyl)ethylamino]-9-(β -D-ribofuranosyl)purine, (S. Kusachi et al, J. Med. Chem., 1985, 28, 1636).

Example 55: 6-(N-Methylphenylamino)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 358 [M+H]⁺.

10 Example 56: 9-(β -D-Ribofuranosyl)-6-(1,2,4,5-tetrahydro-3H-benzazepin-3-yl)purine; mass spectrum m/z 398 [M+H]⁺.

Example 57: 9-(β -D-Ribofuranosyl)-6-(1,2,3,4-tetrahydro-2-isoquinolyl)purine; mass spectrum m/z 384 [M+H]⁺.

15 Example 58: 6-(4-Methylpiperazinyl)-9-(β -D-ribofuranosyl)purine, (H. Vorbrueggen and K. Krolkiewicz, Liebigs Ann. Chem., 1976, 745).

Example 59: 9-(β -D-Ribofuranosyl)-6-(1,3,4,5-tetrahydro-2H-benzazepin-2-yl)purine; mass spectrum m/z 398 [M+H]⁺.

Example 60: 6-[2-(4-Cyanomethylphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 411 [M+H]⁺.

20 Example 61: 6-(2,3-Dihydro-1-indolyl)- 9-(β -D-ribofuranosyl)purine; mass spectrum m/z 370 [M+H]⁺.

Example 62: 9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl)purine; mass spectrum m/z 416 [M+H]⁺.

25 Example 63: 9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl)purine; mass spectrum m/z 400 [M+H]⁺.

Example 64: 6-(8-Aminosulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 477 [M+H]⁺.

- Example 65: 6-[2-(3,4-Dimethoxyphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine, (H. Vorbrueggen and K. Krolikiewicz, Liebigs Ann. Chem., 1976, 745).
- 5 Example 66: 6-[2-(4-Hydroxyphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine, (Shikita et al, Chem. Pharm.Bull., 1974, 22, 1410).
- Example 67: 6-(2-Isoindolinyl)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 370 [M+H]⁺.
- 10 Example 68: 6-(7-Aminosulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl)-9-(β -D-Ribofuranosyl)purine; mass spectrum m/z 477 [M+H]⁺.
- Example 69: 6-(N-Cyclohexylmethylamino)-9-(β -D-ribofuranosyl)purine, (Patent No. DE2148838).
- 15 Example 70: 6-(N-Hexylmethylamino)-9-(β -D-ribofuranosyl)purine, (Patent No. DE2148838).
- Example 71: 6-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-ylamino)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 460 [M+H]⁺.
- 20 Example 72: 6-[N-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-yl)methylamino]-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 474 [M+H]⁺.
- Example 73: 6-[N-(5-Aminopentyl)methylamino]-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 367 [M+H]⁺.
- 25 Example 74: 6-[(5-Chloro-2-methoxyphenyl)methylamino]-9-(β -D-ribofuranosyl)purine, (Patent No. DE2148838).
- Example 75: 6-[(2-Methylphenyl)methylamino]-9-(β -D-ribofuranosyl)purine, (A. M. Aronov et al, J. Med. Chem., 1998, 41, 4790).
- Example 76: 6-(Hexamethyleneimino)-9-(β -D-ribofuranosyl)purine, (H. Vorbrueggen and K. Krolikiewicz, Liebigs Ann. Chem., 1976, 745); mass spectrum (ESI) m/z 350[M+H]⁺.
- 25 Example 77: 6-(1-Pyrrolidinyl)-9-(β -D-ribofuranosyl)purine, (M. Legraverend et al, Tetrahedron, 1984, 40, 709); mass spectrum (ESI) m/z 322 [M+H]⁺.

- Example 78: 6-(4-Hydroxypiperidin-1-yl)- 9-(β -D-ribofuranosyl)purine, (Patent No.DE 2157036); mass spectrum (ESI) m/z 352 [M+H]⁺.
- Example 79: 6-(1-Piperidinyl)-9-(β -D-ribofuranosyl)purine, (M. Legraverend et al, Tetrahedron, 1984, 40, 709); mass spectrum (ESI) m/z 336 [M+H]⁺.
- 5 Example 80: 6-(2-Propenyl)amino-9-(β -D-ribofuranosyl)purine, (M. H. Fleysher et al, J. Med. Chem., 1980, 23, 1448); mass spectrum (ESI) m/z 308 [M+H]⁺.
- Example 81: 6-(2-Propynyl)amino-9-(β -D-ribofuranosyl)purine, (M. H. Fleysher et al, J. Med. Chem., 1980, 23, 1448); mass spectrum (ESI) m/z 306 [M+H]⁺.
- 10 Example 82: 6-(1-Methyl)ethylamino-9-(β -D-ribofuranosyl)purine, (A. M. Aronov et al, J. Med. Chem., 1998, 41, 4790) mass spectrum (ESI) m/z 310 [M+H]⁺.
- Example 83: 6-bis-(2-Propenyl)amino-9-(β -D-ribofuranosyl)purine, (Patent No. DE 2338963); mass spectrum (ESI) m/z 348 [M+H]⁺.
- 15 Example 84: 6-(2-Phenylethyl)methylamino-9-(β -D-ribofuranosyl)purine ,(S. Kusachi et al, J. Med. Chem., 1985, 28, 1636); mass spectrum (ESI) m/z 386 [M+H]⁺.
- Example 85: 6-Ethylmethylamino- 9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 310 [M+H]⁺.
- 20 Example 86: 6-bis-[(3-Methyl)butylamino]-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 408 [M+H]⁺.
- Example 87: 6-(4-Aminophenyl)methylamino-9-(β -D-ribofuranosyl)purine, (M.J.Robins et al, Nucleosides and Nucleotides, 1994, 13, 1627).
- 25 Example 88: 6-(2-Pyridylmethyl)amino-9-(β -D-ribofuranosyl)purine ,(S. Kusachi et al, J. Med. Chem., 1985, 28, 1636); mass spectrum (ESI) m/z 359 [M+H]⁺.
- Example 89: 6-(2-Hydroxyethyl)methylamino-9-(β -D-ribofuranosyl)purine (P.F.Guengerich and V.M.Raney, J.Amer.Chem.Soc., 1992,114,1074).
- Example 90: 6-Dipropylamino-9-(β -D-ribofuranosyl)purine, (M. de Zwart et al, Nucleosides and Nucleotides, 1998, 17, 969).

Example 91

Starting with 2',3',5'-tris-O-(tert-butyldimethylsilyl)adenosine in manner analogous to that described by K. Aritomo, T. Wada and M. Sekine, J. Chem. Soc. Perkin Trans.1, 1995,1837 was prepared 6-[2-phenyl-(N-propionyl)ethylamine]-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 428 [M+H]⁺.

Example 92

Starting with 2',3',5'-tris-O-(tert-butyldimethylsilyl)adenosine in manner analogous to that described by K. Aritomo, T. Wada and M. Sekine , J. Chem. Soc. Perkin Trans.1, 1995,1837 was prepared 6-(N-benzoyl-2-phenylethylamine)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 476 [M+H]⁺.

Example 93

Starting with adenosine in manner analogous to that described by T. Itaya et al, Chem. Pharm. Bull., 1977, 25, 1449 was prepared 1-benzyl-6-imino-9-(β -D-ribofuranosyl)purine.

Example 94

Starting with 6-(2-phenylethylamino)-9-(β -D-ribofuranosyl)purine (prepared in a manner analogous to that described in example 83) and in manner analogous to that described by T. Itaya et al, Chem. Pharm. Bull., 1977, 25, 1449 was prepared 1-methyl-6-(2-phenylethylamino)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 386 [M+H]⁺.

25

Example 95

A solution of 0.34g of 2-amino-6-chloro-9-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)purine in 5ml of a 2M solution of methylamine in methanol was heated under nitrogen under reflux overnight. The solvents were removed by

evaporation and the residue purified by preparative HPLC to give 10mg of 2-amino-6-methylamino-9-(β -L-ribofuranosyl)purine as a pale yellow solid; mass spectrum (ESI) m/z 297[M+H]⁺.

5 The 2-amino-6-chloro-9-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)purine used as the starting material was prepared as follows:

A suspension of 38mg of 2-amino-6-chloropurine in 1ml of anhydrous acetonitrile was treated with 0.22ml of bis(trimethylsilyl)acetamide and heated at reflux for 15 min. To the resulting solution was added a solution of 95mg of 1-O-acetyl-2,3,5-tri-O-benzoyl-L-ribose in 1ml of anhydrous acetonitrile followed by 51 μ l of trimethylsilyl trifluoromethanesulphonate. The solution was heated at reflux under nitrogen for 2.5 hours. After cooling to room temperature the solution was evaporated and the residue dissolved in dichloromethane and washed twice with water. The solution was dried over anhydrous magnesium sulphate, filtered and evaporated to give crude 2-amino-6-chloro-9-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)purine which was used without further purification; mass spectrum (ESI) m/z 614 [M+H]⁺.
10
15

Example 96

Reaction of 2-amino-6-chloropurine with 1-O-acetyl-2,3,5-tri-O-benzoyl-D-ribose followed by treatment of the intermediate 2-amino-6-chloro-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine with methylamine in methanol in an analogous manner to that described in example 95 gave 2-amino-6-methylamino-9-(β -D-ribofuranosyl)purine (R. Saladino et al, Tetrahedron, 1996, 52, 6759); mass spectrum (ESI) m/z 297[M+H]⁺.
20
25

Example 97

Reaction of 2-amino-6-chloropurine with 1-O-acetyl-2,3,5-tri-O-benzoyl-D-ribose followed by treatment of the intermediate 2-amino-6-chloro-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine with morpholine in methanol in an analogous manner to that described in example 95 gave 2-amino-6-(4-morpholinyl)-9-(β -D-ribofuranosyl)purine (H.Vorbrueggen and K.Krolikiewicz, Justus Leibigs Ann.Chem.,1976, 745).
30

Example 98

Reaction of 2-amino-6-chloropurine with 1-O-acetyl-2,3,5-tri-O-benzoyl-D-ribose followed by treatment of the intermediate 2-amino-6-chloro-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine with pyrrolidine in methanol in an analogous manner to that described in example 95 gave 2-amino-6-(1-pyrrolidinyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 337 [M+H]⁺.

Example 99

A suspension of 84mg of 2,4-diaminopurine in 2ml of anhydrous acetonitrile was treated with 0.55ml of bis(trimethylsilyl)acetamide and the solution heated at reflux for 15 min to give a solution. To the solution was added a solution of 237mg of 1-O-acetyl-2,3,5-tri-O-benzoyl-L-ribose in 2ml of anhydrous acetonitrile. The solution was heated at reflux under nitrogen for 16 hours. After cooling to room temperature the solution was evaporated and the residue dissolved in dichloromethane and washed with water. The dichloromethane solution was dried over anhydrous magnesium sulphate, filtered and evaporated. The residue was dissolved in 10ml of a 2M solution of ammonia in methanol and the solution stirred at room temperature for 42 hours then evaporated. The residue was purified by preparative HPLC to give 50mg of 2,6-diamino-9-(β -L-ribofuranosyl)purine, (D.M.Brown et al, Nucleosides and Nucleotides, 1999, 18, 2521); mass spectrum (ESI) m/z 283[M+H]⁺.

Example 100

Reaction of 2,6-diaminopurine with 1-O-acetyl-2,3,5-tri-O-benzoyl-D-ribose followed by treatment of the intermediate 2,6-diamino-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine with ammonia in methanol in an analogous manner to that described in example 99 gave 2,6-diamino-9-(β -D-ribofuranosyl)purine (also available commercially from ICN Biomedicals Inc.).

- 139 -

Example 101

A mixture of 4.5 g of 2,6-dichloro-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine, 1.1g of pyrrolidine and 2.8ml of triethylamine in 50ml of benzene was stood at room temperature for 1 hour then washed with water, dried and evaporated. The residue was dissolved in a saturated solution of ammonia in methanol and the solution stood overnight at room temperature. The solution was evaporated and the residue recrystallised from n-butanol to give 2.5g of 2-chloro-6-(1-pyrrolidinyl)-9-(β -D-ribofuranosyl)purine (W. Kampe et al, Patent No. DE 2157036) of melting point 229°C; mass spectrum (ESI) m/z 356 [M+H]⁺.

10

Example 102

By an analogous procedure to that described in example 101 starting with 2,6-dichloro-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine and hexamethyleneimine was prepared 2-chloro-6-(1-hexamethyleneimino)-9-(β -D-ribofuranosyl)purine, (W. Kampe et al, Patent No. DE 2157036); mass spectrum (ESI) m/z 384 [M+H]⁺.

15

Example 103

By an analogous procedure to that described in example 101 starting with 2,6-dichloro-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine and 4-hydroxypiperidine was prepared 2-chloro-6-(4-hydroxy-1-piperidinyl)-9-(β -D-ribofuranosyl)purine (W. Kampe et al, Patent No. DE 2157036); mass spectrum (ESI) m/z 386 [M+H]⁺.

20
25

Example 104

By a procedure analogous to that described by Kissman et al, J. Amer. Chem. Soc., 1955, 77,18 was prepared 6-[(N-cyclohexyl)methylamino]-2-methylthio-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 410 [M+H]⁺.

- 140 -

Example 105

A solution of 30g of adenosine and 16.4ml of 2,5-dimethoxytetrahydrofuran in 70ml of glacial acetic acid was heated at reflux temperature for 1 hour. After cooling to room temperature the mixture was concentrated under reduced pressure, and the residual oil triturated with acetone, filtered and the filtrate evaporated. The residue was purified by column chromatography on silica gel using methanol/dichloromethane (5:95) as the eluent to give 17.0g of 6-(1-pyrrolyl)-9-(β -D-ribofuranosyl)purine as a light orange solid; mass spectrum (ESI) m/z 318 [M+H].

10

Example 106

Reaction of 6-amino-9-(β -D-arabinofuranosyl)purine with dimethoxytetrahydrofuran in an analogous manner to that described in example 105 gave 6-(1-pyrrolyl)-9-(β -D-arabinofuranosyl)purine as a light brown solid of melting point 212-213°C; mass spectrum (ESI) 318 [M+H]⁺.

15

Example 107

A solution containing 150mg of 6-amino-9-(β -D-ribofuranosyl)purin-8-(7H)-one and 74mg of 2,5-dimethoxytetrahydrofuran in 5ml glacial acetic acid was heated under nitrogen at 110°C for 1 hour. The solvents were then evaporated under low vacuum to give a brown residue, which was purified by flash chromatography on silica-gel using methanol/dichloromethane (1:9) for the elution to give 18mg of 6-(1-pyrrolyl)-9-(β -D-ribofuranosyl)purin-8(7H)-one as a white solid; mass spectrum (ESI) m/z 334 [M+H]⁺.

20

25

Example 108

A solution containing 150mg of 9-(3'-deoxy- β -D-ribofuranosyl)adenosine and 83mg of 2,5-dimethoxytetrahydrofuran in 5ml glacial acetic acid was heated under nitrogen at 110°C for 2 hours. The solvents were then evaporated under low vacuum to give a beige solid which was purified by flash chromatography on silica-

30

gel using methanol/dichloromethane (1:49) for the elution to give 70mg of 9-(3-deoxy- β -D-ribofuranosyl)-6-(1-pyrrolyl) purine as a white solid of melting point 175-176°C; mass spectrum (ESI) m/z 302 [M+H]⁺.

5

Example 109

A solution of 0.51g of 6-(1-pyrrolyl)-9-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)purine and 20ml of a 33% aqueous ammonia solution in 30ml of methanol/tetrahydrofuran (1:1), was heated at 50° C for 2 hours. After cooling to room temperature the mixture was evaporated, diluted with 50ml of water and extracted twice with 50ml diethyl ether followed by 50ml ethyl acetate. The combined organic extracts were dried over anhydrous sodium sulphate, concentrated under reduced pressure and the mixture purified by column chromatography on silica gel using methanol/dichloromethane (5:95) as the eluent, to give 0.12g of 6-(1-pyrrolyl)-9-(β -L-ribofuranosyl)purine as a white solid of melting point 114-115°C; mass spectrum (ESI) 318 [M+H]⁺.

The 6-(1-pyrrolyl)-9-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)purine used as a starting material was prepared as follows:

To a suspension of 1.0g of 6-(1-pyrrolyl)purine (prepared according to K.G.Estep et al, J.Med.Chem., 1995, 38, 2582) and 0.97g of 1-O-acetyl-2,3,5-tri-O-benzoyl- β -L-ribofuranose in 30ml of 1,2-dichloroethane was added dropwise 2.30g of N-methyl-N-trimethylsilyl trifluoroacetamide, and the mixture heated to 80 ° C. Following addition of 0.635g of trimethylsilyl trifluoromethane sulphonate, dropwise, the mixture was stirred at 80°C overnight. After cooling to room temperature, the mixture was diluted with 60ml of dichloromethane and washed four times with a saturated solution of aqueous sodium hydrogen carbonate. The organic extract was dried over sodium sulphate, filtered and evaporated and the residue purified by flash column chromatography on silica gel using ethyl acetate/hexane (10:90) for the elution to give 0.56g of 6-(1-pyrrolyl)-9-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)purine as a white solid; mass spectrum (ESI) 630 [M+H].

Example 110

Reaction of 6-(1-indolyl)purine (M. Haidoune and R Mornet, J. Heterocyclic Chem., 1994, 31, 1461) with 1-O-acetyl-2,3,5-tri-O-benzoyl- β -D-ribofuranose followed deprotection in an analogous manner to that described in example 109 gave 6-(1-indolyl)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 368 [M+H]⁺.

Example 111

Reaction of 6-(1-imidazol-yl)purine (G. E. Estep et al, J. Med. Chem., 1995, 38, 2582) with 1-O-acetyl-2,3,5-tri-O-benzoyl- β -D-ribofuranose followed deprotection in an analogous manner to that described in example 109 gave 6-(1-imidazolyl)-9-(β -D-ribofuranosyl)purine; mass spectrum m/z 319 [M+H]⁺.

Example 112

150 μ l of a 1M solution of sodium methoxide in methanol was added to a stirring solution of 0.445g of 6-(1,2,4-triazol-1-yl)-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine in 10ml of anhydrous methanol. After stirring overnight at room temperature a few drops of glacial acetic acid were added and the mixture concentrated under reduced pressure. The mixture was purified by column chromatography on silica gel using an eluent of methanol/dichloromethane (10:90) to give 0.2g of 9-(β -D-ribofuranosyl)-6-(1,2,4-triazol-1-yl)purine as a white solid, mass spectrum (ESI) 320 [M+H].

The 6-(1,2,4-triazol-1-yl)-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine used as a starting material was prepared as follows:

3.7ml of Phosphorous oxychloride followed by 30ml of triethylamine were added dropwise to a solution of 13.1g of 1,2,4-triazole in 150ml of acetonitrile at <5 ° C. After stirring for 1 hour, a suspension of 5.0g of 2',3',5'-tri-O-acetylinosine in 150ml of acetonitrile was added, and the mixture stirred at room temperature overnight. The mixture was filtered, diluted with 100ml of ethyl acetate and extracted twice with 100ml of a saturated solution of aqueous sodium hydrogen carbonate. The organic extract was dried over anhydrous sodium sulphate and concentrated under reduced pressure. The mixture was purified by column

chromatography on silica gel using methanol/dichloromethane (5:95) for the elution to give 2.7g of 6-(1,2,4-triazol-1-yl)-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine as a white foam, mass spectrum (ESI) 446 [M+H].

5

Example 113

Reaction of 6-(1-pyrazolyl)-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine with sodium methoxide in an analogous manner to that described in example 112 followed by purification by supercritical fluid chromatography gave 6-(1-pyrazolyl)-9-(β -D-ribofuranosyl)purine as a white solid, mass spectrum (ESI) 319 [M+H].

10 The 6-(1-pyrazolyl)-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine used as a starting material was prepared as follows:

15 0.78ml of chlorotrimethylsilane was added dropwise to a stirring solution of 0.372g of 6-(1-pyrazolyl)purine (prepared according to K.G.Estep et al, J.Med.Chem., 1995, 38, 2582) 1.0g of β -D-ribofuranose-1-acetate-2,3,5-tribenzoate, 1.62g of nonafluoro-1-butanesulfonic acid and 0.3ml of hexamethyldisilazane in 30ml of acetonitrile, and the mixture heated at reflux temperature for 21 hours. After cooling to room temperature, the mixture was diluted with 30ml of dichloromethane and washed with 50ml of a saturated aqueous solution of sodium hydrogen carbonate. The organic extract was dried over anhydrous sodium sulphate and concentrated under reduced pressure. The mixture was purified by column chromatography on silica gel using an eluent of methanol/dichloromethane (5:95) to give 0.06g of 6-(pyrazol-1-yl)-9-(2,3,5-tri-O-benzoyl- β -D-ribofuranosyl)purine as a yellow solid, mass spectrum (ESI) 630 [M+H].

25

Example 114

30 By the procedure of V. Samano, R. W. Robins and M. J. Robins, J. Amer. Chem. Soc., 1994, 116, 9331 was prepared 9-(β -D-ribofuranosyl) 6-(1,2,4-triazol-4-yl)purine; mass spectrum (ESI) m/z 320[M+H]⁺.

- 144 -

Example 115

By a procedure analogous to that of J. A. Montogomery, J. A. Secrist and C. A. Krauth, US Patent No. 5,102,873 starting with adenosine was prepared 6-(2-phenylethylamino)- 9-(β -D-ribofuranosyl)purine-1-oxide.

5

Example 116

By the procedure of Yamazaki et al, Chem. Pharm. Bull., 1968, 16, 2172 was prepared 6-methylamino-9-(β -D-ribofuranosyl)purin-2(1H)-one of melting point 270°C (decomposition).

10

Example 117

By the procedure of G.R.Gough and H.M.Maguire, J.Med.Chem., 1967,10, 475 was prepared 2-methoxy-6-methylamino-9-(1- β -D-ribofuranosyl)purine of melting point 142°C (decomposition).

15

Example 118

By the procedure of T. Schaeffer, J. Amer. Chem. Soc., 1958, 80,3738 starting with 2-chloroadenosine (Aldrich Chemical Co.) was prepared 2-methoxyadenosine.

20

Example 119

By the procedure of J. F. Gerster and R. K. Robins, J. Org. Chem., 1966, 31, 3528 was prepared 2-amino-6-chloro-9-(β -D-ribofuranosyl)purine.(Sigma-Aldrich Chemical Co.).

25

Example 120

By the procedure of Johnson et al, J.Amer.Chem.Soc., 1958, 80; 699 starting with 6-chloro-9-(β -D-ribofuranosyl)purine was prepared 6-methoxy-9-(1- β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 283 [M+H]⁺.

5

Example 121

By the procedure of C. W. Noell and R. K. Robins, J. Med. Pharm. Chem., 1962, 5, 1074 was prepared 2-amino-6-benzylthio-9-(β -D-ribofuranosyl)purine.

10

Example 122

By the procedure of W. Kampe et al, Patent No. ZA 6707630 was prepared 6-benzylthio-2-hydroxy-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 391[M+H]⁺.

15

Example 123

By the procedure of B. S. Schultz and W. Pleiderer, Tet. Lett., 1985, 26, 5421 from guanosine was prepared 9-(β -D-ribofuranosyl)purine-2,6,8(1H,3H,7H)-trione; mass spectrum (ESI) m/z 342[M+CH₃CN+H]⁺.

20

Example 124

By the procedure of C. B. Reese and R. Saffhill, J. Chem. Soc. Perkin Trans. 1, 1972, 2937 was prepared 2-(acetylamino)inosine; mass spectrum (ESI) m/z 326[M+H]⁺.

Example 125

A mixture of 0.5g of 8-bromo-adenosine and 0.5ml of water was treated with 1ml of a 33% solution of methylamine in ethanol. The mixture was heated at 70°C for 12 hours then evaporated to dryness. The crude product (0.54g) was purified by flash column chromatography on silica gel using methanol/dichloromethane (1:9 to 3:9) for the elution to give 0.34g of 8-(methylamino)adenosine (J. B. Chattopadhyaya and C. B. Reese, Synthesis, 1977, 725) as a white solid of melting point >250°C; mass spectrum (ESI) m/z 297 [M+H]⁺.

In a manner analogous to that described in example 125 starting with 8-bromo-adenosine and the appropriate amine in ethanol or aqueous ethanol were prepared the following examples:

Example 126: 8-(2-Phenylethylamino)adenosine.

Example 127: 8-Benzylamino-adenosine (A.M.Aronov and M.H.Gelb, Biorg.and Med.Chem.Lett., 1998,24,3505) of melting point 213-216°C.

Example 128: 8-(1-Piperidinyl)adenosine (A.M.Aronov and M.H.Gelb, Biorg.and Med.Chem.Lett. 1998,24,3505) of melting point 207-209°C (decomposition).

Example 129: 8-(Dimethylamino)adenosine (A.M.Aronov and M.H.Gelb, Biorg.and Med.Chem.Lett. 1998,24,3505) of melting point 205-207°C.

Example 130: 8-(3-Phenylpropylamino)adenosine of melting point 180-183°C.

Example 131: 8-(4-Morpholinyl)adenosine of melting point 210-213°C.

Example 132: 8-(N-Methyl-2-phenylethylamino)adenosine of melting point 118-120°C.

Example 133: 8-(3-Pyridylmethylamino)adenosine of melting point 235-237°C (decomposition).

Example 134: 8-(Ethylamino)adenosine (R.A.Long and R.K.Robins, J.Org.Chem., 1967, 32, 2751) of melting point 260-170°C.

- Example 135: 8-(1,2,3,4-Tetrahydro-2-isoquinolyl)adenosine of melting point 145-150°C (decomposition).
- Example 136: 8-[2-(4-Morpholinyl)ethylamino]adenosine of melting point 210-215°C.
- 5 Example 137: 8-(Hexylamino)adenosine (Patent No. JP53124293) of melting point 209-212°C.
- Example 138: 8-(2-Cyclohexylethylamino)adenosine of melting point 203-205°C.
- Example 139: 8-(2(R,S)-Phenylpropylamino)adenosine of melting point 159-161°C (decomposition).
- 10 Example 140: 8-[2-(4-Methylphenyl) ethylamino]adenosine of melting point 117-124°C (decomposition).
- Example 141: 8-[2-(1-Methyl-2-pyrrolyl) ethylamino]adenosine of melting point 225-228°C.
- 15 Example 142: 8-[2-(4-Aminosulphonylphenyl)ethylamino]adenosine of melting point 157-163°C (decomposition).
- Example 143: 8-(4-Phenyl-1-piperazinyl)adenosine of melting point 220-223°C (decomposition).
- 20 Example 144: 8-(2-(4-Imidazolyl)adenosine (T. Prakash and K.N.Ganesh, J.Chem.Soc.Chem.Commun.,1994,1357) of melting point 148-156°C (decomposition).
- Example 145: 8-(1-Naphthylmethylamino)adenosine of melting point 140-150°C.
- Example 146: 8-[2-(4-Hydroxyphenyl)ethylamino]adenosine of melting point 262-265°C (decomposition).
- Example 147: 8-(4-Phenylbutylamino)adenosine of melting point 190°C.
- 25 Example 148: 8-[2-(4-Chlorophenyl)ethylamino]adenosine of melting point 155-158°C (decomposition).
- Example 149: 8-[2-(2,4-Dichlorophenyl)ethylamino]adenosine of melting point 164-168°C (decomposition).

Example 150: 8-(2-Propenylamino)adenosine of melting point 234-237°C (decomposition). Example 163: 8-[(4-tert-Butyl)benzylamino]adenosine of melting point 187-190°C.

Example 164: 8-(1(R)-Phenylethylamino)adenosine of melting point 120-130°C.

5 Example 165: 8-(1(S)-Phenylethylamino)adenosine of melting point 112-130°C.

Example 166: 8-(6-Phenylhexylamino)adenosine of melting point 165-167°C.

Example 167: 8-[2-Hydroxy-1(S)-phenyl]ethylamino]adenosine of melting point 110-125°C.

10 By a procedure analogous to that described in example 125 from 8-bromo-2'-deoxyadenosine were prepared the following examples:

Example 168: 2'-Deoxy-8-(2-phenylethylamino)adenosine of melting point 192-195°C.

15 Example 169: 2'-Deoxy-8-(3-phenylpropylamino)adenosine of melting point 198-201°C.

Example 170: 8-Benzylamino-2'-deoxyadenosine of melting point 132-134°C.

Example 171: 2'-Deoxy-8-(4-phenylbutylamino)adenosine of melting point 168-171°C.

20 Example 172: 2'-Deoxy-8-(6-phenylhexylamino)adenosine of melting point 159-161°C.

Example 173

By a procedure analogous to that described in example 125 from 8-bromoinosine was prepared 8-(4-morpholinyl)inosine (M. Sechenova, Fiziol.Zh.SSSR, 1989, 75, 457).

Example 174

By a procedure analogous to that described in example 125 from 8-bromoinosine was prepared 8-benzylaminoinosine (Chattopadhyaya and Reese, Synthesis, 1978, 908) of melting point 225-228°C.

Example 175

By the procedure of G. S. Buenger, Synthesis, 1990, 962 starting with 8-bromoadenosine was prepared 8-(methylthio)adenosine of melting point 254-255°C.

Example 176

By an analogous procedure to that of G. S. Buenger, Synthesis, 1990, 962 starting with 8-bromoadenosine was prepared 8-(benzylthio)adenosine (E, Liepins et al, Bioorg. Khim., 1988, 14, 1393) of melting point 206-210°C.

Example 177

By the procedure of G. S. Buenger, Synthesis, 1990, 962 starting with 8-bromoadenosine was prepared 8-(benzyloxy)adenosine of melting point 199-201°C.

Example 178

By an analogous procedure to that of G. S. Buenger, Synthesis, 1990, 962 starting with 8-bromoadenosine was prepared 8-ethoxyadenosine of melting point 172-175°C.

- 150 -

Example 179

By the procedure of Holmes and Robins, J. Amer. Chem. Soc., 1964, 86, 1242 starting with 8-bromoadenosine was prepared 6-amino-9-(β -D-ribofuranosyl)purine-8(7H)-thione of melting point 242-248°C (decomposition).

5

Example 180

By the procedure of H. Steinmaus et al, J. Org. Chem., 1971, 36, 3594 starting with adenosine was prepared 8-[(1-hydroxy-1-methyl)ethyl]adenosine.

10

Example 181

A solution of 0.31g of 9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)-6-(3-thienyl)purine in 3ml of anhydrous methanol was treated with 67 μ l of a 1M solution of sodium methoxide in methanol. The mixture was stirred at room temperature for 2 hours during which time a white precipitate separated. A few drops of glacial acetic acid were added and the mixture was evaporated to dryness under reduced pressure. Recrystallisation of the residue from ethanol gave 0.11g of 9-(β -D-ribofuranosyl)-6-(3-thienyl)purine as a white solid of melting point 166-167°C (decomposition); mass spectrum (ESI) m/z 335[M+H]⁺.

15

The 9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)-6-(3-thienyl)purine used as the starting material was prepared as follows:

20

A mixture containing 0.5g of 9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)-6-chloropurine, 0.23g of thiophene-3-boronic acid, 0.21g of anhydrous potassium carbonate and 0.034g of tetrakis-(triphenylphosphine)palladium in 24ml of anhydrous toluene was stirred under nitrogen and heated at 100°C for 5 hours. After cooling the mixture was diluted with 50ml of ethyl acetate and washed with 20ml of water and 20ml of brine. The solution was dried over anhydrous magnesium sulphate, filtered and evaporated to yield a gum. This was purified by flash chromatography on silica gel using ethyl acetate/hexane(1:1) for the elution to give 0.31g of 9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)-6-(3-thienyl)purine as a gum; mass spectrum (ESI) m/z 461[M+H]⁺.

25

30

Example 182

Reaction of 6-chloro-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine with phenylboronic acid followed by deprotection in an analogous manner to that described in example 181 gave 6-phenyl-9-(β -D-ribofuranosyl) purine, (M. Hocek, A. Holy, I. Votruba and H. Dvorakova, J. Med. Chem., 2000, 43, 1817) as a white solid of melting point 224-225°C; mass spectrum (ESI) m/z 329[M+H]⁺.

Reaction of 50mg samples of 6-chloro-9-(tri-O-acetyl- β -D-ribofuranosyl)purine with a range of arylboronic acids in an analogous manner to that described in example 181 was carried out in parallel using a Mettler Toledo Myriad reactor. The intermediate crude 6-aryl-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purines were purified using a Jones Flashmaster II sequential chromatography system using ethyl acetate/hexane for the elution before deprotection using sodium methoxide in methanol in an analogous manner to that described in example 181 to give the 6-aryl-9-(β -D-ribofuranosyl)purines listed below:

Example 183: 6-(4-Fluorophenyl)-9-(β -D-ribofuranosyl)purine (M Hocek et al, J Med Chem, 2000, 43, 1817); mass spectrum (ESI) m/z 347[M+H]⁺.

Example 184: 6-(4-Chlorophenyl)-9-(β -D-ribofuranosyl)purine (M Hocek et al, J Med Chem, 2000, 43, 1817); mass spectrum (ESI) m/z 363[M+H]⁺.

Example 185: 6-(4-Methylphenyl)-9-(β -D-ribofuranosyl)purine (M Hocek et al, J Med Chem, 2000, 43, 1817); mass spectrum (ESI) m/z 343[M+H]⁺.

Example 186: 6-(4-Methoxyphenyl)-9-(β -D-ribofuranosyl)purine (M Hocek et al, J Med Chem, 2000, 43, 1817); mass spectrum (ESI) m/z 359[M+H]⁺.

Example 187: 9-(β -D-Ribofuranosyl)-6-(1-thianthrenyl)purine; mass spectrum (ESI) m/z 467[M+H]⁺.

Example 188: 6-(4-Biphenylyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 405[M+H]⁺.

Example 189: 6-(4-Methylthiophenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 375[M+H]⁺.

Example 190: 6-(2-Methylphenyl)-9-(β -D-ribofuranosyl)purine (M Hocek et al, J Med Chem, 2000, 43, 1817); mass spectrum (ESI) m/z 343[M+H]⁺.

Example 191: 6-(9-Phenanthrenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 429[M+H]⁺.

5 Example 192: 9-(β -D-Ribofuranosyl)-6-(3-trifluoromethylphenyl)purine; mass spectrum (ESI) m/z 397[M+H]⁺.

Example 193: 6-(2-Phenoxyphenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 421[M+H]⁺.

10 Example 194: 6-(4-tert-Butylphenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 385[M+H]⁺.

Example 195: 9-(β -D-Ribofuranosyl)-6-(2-trifluoromethoxyphenyl)purine; mass spectrum (ESI) m/z 413[M+H]⁺.

Example 196: 6-(4-Phenoxyphenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 421[M+H]⁺.

15 Example 197: 6-(3-Methoxyphenyl)-9-(β -D-ribofuranosyl)purine (M Hocek et al, J Med Chem, 2000, 43, 1817); mass spectrum (ESI) m/z 359[M+H]⁺.

Example 198: 6-(2-Naphthyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 379[M+H]⁺.

20 Example 199: 6-(3-Biphenylyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 405[M+H]⁺.

Example 200: 6-[4-(2-Methylpropyl)phenyl]-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 385[M+H]⁺.

Example 201: 6-(3-Fluorophenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 347[M+H]⁺.

25 Example 202: 9-(β -D-Ribofuranosyl)-6-(4-trifluoromethylphenyl)purine; mass spectrum (ESI) m/z 397[M+H]⁺.

Example 203: 9-(β -D-Ribofuranosyl)-6-(4-trifluoromethylphenyl)purine; mass spectrum (ESI) m/z 373[M+H]⁺.

Example 204: 6-[3-(1-methyl)ethylphenyl]-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 371[M+H]⁺.

Example 205: 9-(β -D-Ribofuranosyl)-6-(4-trifluoromethoxyphenyl)purine; mass spectrum (ESI) m/z 413[M+H]⁺.

- 5 Example 206: 6-(4-Ethylphenyl)-9-(β -D-ribofuranosyl)purine; mass spectrum (ESI) m/z 357[M+H]⁺.

Example 207

Reaction of 2-amino-6-chloro-9-(2,3,5-tri-O-acetyl- β -D-ribofuranosyl)purine with phenylboronic acid followed by deprotection in an analogous manner to that described in example 181 gave 2-amino-6-phenyl-9-(β -D-ribofuranosyl)purine (M Hocek, A. Holy, I. Votruba and H. Dvorakova, J. Med. Chem., 2000, 43, 1817) as a white solid of melting point 187-190°C; mass spectrum (ESI) m/z 344[M+H]⁺.

15

Example 208

A solution of 0.2g of 2'3'5'-tri-O-benzoyl-5-ethyluridine in 1ml of anhydrous methanol was treated with 0.05ml of 1M sodium methoxide solution in methanol. The solution was stirred at room temperature for 2 hours. A few drops of glacial acetic acid was added and the mixture evaporated to dryness. The solid residue was purified by flash column chromatography on silica gel using ethyl acetate/isohexane for the elution to give 50mg of 5-ethyluridine (C. Nakayama et al, J. Carbohydr. Nucleosides and Nucleotides, 1979, 6, 295) of melting point 180-181°C; mass spectrum (ESI) 273[M+H]⁺.

25 The 2'3'5'-tri-O-benzoyl-5-ethyluridine used as the starting material was prepared as follows:

A mixture of 0.84g of 5-ethyluracil, 2mg of ammonium sulphate and 3.9ml of hexamethyldisilazane was stirred under nitrogen and heated under reflux for 3.5 hours to give a clear solution. The solution was evaporated under reduced pressure to give an oil which was dissolved in 5ml of anhydrous acetonitrile. This solution

was added to a solution of 3.0g of 1-O-acetyl-2,3,5-tri-O-benzoyl-D-ribose in 20ml of anhydrous acetonitrile. The mixture was cooled in ice at <5°C and treated with 1.4ml of stannic chloride in three portions during 5 min then stirred at room temperature overnight. The mixture was treated with 12ml of water and adjusted to pH 8 by addition of solid sodium bicarbonate. The resulting slurry was filtered through a pad of Hyflo and the filtered solid washed three times with dichloromethane. The combined filtrates were transferred to a separating funnel and the layers separated. The dichloromethane solution was dried over anhydrous sodium sulphate, filtered and evaporated to give 3.3g of white solid residue. This
5 was purified by flash column chromatography on silica gel using ethyl acetate/isohexane (1:1) for the elution to give 2.7g of 2'3'5'-tri-O-benzoyl-5-ethyluridine as a white solid; mass spectrum (ESI) m/z 585[M+H]⁺.
10

In an analogous manner to that described in example 208 were prepared the
15 following examples:

Example 209: 5-[(1-Methyl)ethyl]uridine (B.H.A.Knoblauch et al,
Eur.J.Med.Chem.,1999, 34, 809).

Example 210: 5-Methoxymethyluridine (Patent No. JP57018696).

20 Example 211: 5-Ethoxymethyluridine .

Example 212: 5-Chlorouridine (J.Asakura and M.J.Robins, J.Org.Chem., 1990, 55, 4928).

Example 213: 5-Methyl-1-(β -L-ribofuranosyl)uracil (A.Holy and F.Sorm, Collect.
Czech. Chem. Commun., 1969, 34, 3383; mass spectrum (ESI) m/z 259[M+H]⁺.

25

Example 214

By the procedure of Nakayama et al, J. Carbohydr., Nucleosides,
Nucleotides, 1979, 6, 295 was prepared 1-(β -D-arabinofuranosyl)-5-ethyluracil of
melting point 164-165°C.

- 155 -

Example 215

A solution of 3.0g of 1-(β -D-arabinofuranosyl)uracil and 3.0g of N-bromosuccinimide in 20ml of N,N-dimethylformamide was stirred at room temperature for 1 hour. The solution was evaporated to dryness and the residual yellow oil stirred with a mixture of ethanol and chloroform (4:1) until a fine solid crystallised. After cooling the solid was filtered off washed with ethanol and diethyl ether and dried to give 2.3g of 1-(β -D-arabinofuranosyl)-5-bromouracil, (R.F.Shinazi et al, J. Med. Chem.,1979, 22, 1273). Recrystallisation from ethanol gave analytically pure material with melting point 227°C (decomposition).

10

Example 216

By the procedure of K. Felczak, et al, Nucleosides and Nucleotides, 1993, 12, 245 was prepared 5-methyl-4-thiouridine.

15

Example 217

By the procedure of A. Miah et al., Nucleosides and Nucleotides, 1997, 16, 53 was prepared 4-methoxy-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one.

20

Example 218

By the procedure of K.H.Scheit, Tet. Lett., 1967, 113 was prepared 4-(methylthio)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one; mass spectrum (ESI) m/z 275 [M+H]⁺.

25

Example 219

By an analogous procedure to that of K.H.Scheit, Tet. Lett., 1967, 113 was prepared 5-fluoro-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one; mass spectrum (ESI) m/z 293 [M+H]⁺.

Example 220

By an analogous procedure to that of K.H.Scheit, Tet. Lett., 1967, 113 was prepared 5-methyl-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one; mass spectrum m/z 289 [M+H]⁺.

5

Example 221

By a procedure analogous to that of Fox et al., Tet. Lett. 1966, 4927 was prepared 5-fluoro-4-thiouridine.

10

Example 222

By the procedure of Hoffer et al., J.Amer.Chem.Soc., 1959, 81, 4112 was prepared 1-(2-deoxy- α -D-erythro-pentofuranosyl)-5-fluorouracil.

Example 223

15

By the procedure of Zemlicka et al., J.Amer.Chem.Soc., 1972, 94, 3213 was prepared 2'-Deoxy-5-fluoro-3-methyluridine.

Example 224

20

By an analogous procedure to that of Zemlicka et al., J.Amer.Chem.Soc., 1972, 94, 3213 was prepared 1-(α -D-erythro-2-deoxypentofuranosyl)-5-fluoro-3-methyluracil, (D.J.Adams and G.W.Gooday, Mach. Naturwiss.Tech., 1983, 39).

Example 225

25

A stirred slurry of 1.0g of O₂,2'-anhydrouridine in 22ml of anhydrous chloroform was saturated with hydrogen chloride gas for 5 hours. The solid was filtered off dried and suspended in 150ml of 1,4-dioxane. The suspension was

heated at 75°C under nitrogen until a solution was obtained. After cooling this was evaporated and the residual syrup triturated with 50ml of boiling ethyl acetate. A solid formed which was broken up. After cooling the product was filtered to give 1.05g of 2'-chloro-2'-deoxyuridine (Tetrahedron 1977, 33, 2131). Recrystallisation from ethanol gave analytically pure material of melting point 206-207°C.

The O-2,2'-anhydouridine used as the starting material was prepared as follows:

A mixture of 10.0g of uridine, 11.4g of diphenyl carbonate, 0.2g of sodium hydrogen carbonate and 20ml of N,N-dimethylformamide was stirred under nitrogen and heated at 155°C for 30 min. The solution was cooled and added dropwise to 200ml of anhydrous diethyl ether. After stirring the mixture overnight the precipitated solid was filtered off and washed with methanol and dried to give 6.3g of O2,2'-anhydouridine of melting point 241-244°C.

15

Example 226

A saturated solution of hydrogen bromide in 30ml of trifluoroacetic acid was treated with 1.0 g of O2,2'-anhydouridine. The mixture was stirred for 4 days at room temperature in a sealed flask. The resulting solution was evaporated to dryness to yield a brown syrup which crystallised on standing. Recrystallisation from ethanol gave 2'-bromo-2'-deoxyuridine (Codington et al, J. Org. Chem., 1964, 29, 558) of melting point 194-195°C.

Example 227

By the procedure of J. J. Fox and N. C. Miller, J. Org. Chem., 1963, 28, 936 was prepared 1-(2-deoxy- β -D-lyxofuranosyl)-5-methyluracil of melting point 170-171°C.

Example 228

By the procedure of Johansson et al., Patent No. 5506215 was prepared 3'-deoxy-3'-fluoro-5-methyluridine.

Example 229

A suspension of 2.0g of 2'-deoxy-5-ethyl-5'-O-triphenylmethyluridine in 20ml of benzene and 6.5ml of 1,4-dioxane was stirred and treated with 0.5ml of iodomethane and 0.45g of powdered potassium hydroxide. The mixture was 5 stirred and heated at 40°C for 5 hours then evaporated and the residue dissolved in 2ml of methanol and poured into 100ml of water. The resulting white emulsion was extracted with four 100ml portions of chloroform. The extracts were dried, filtered and evaporated and the residue redissolved in 20ml of 80% acetic acid. The solution was heated at 100°C for 1 hour then evaporated to dryness. The residue 10 was purified by flash column chromatography on silica gel using ethyl acetate for the elution to give 0.25g of 2',3'-dideoxy-5-ethyl-3'-methoxyuridine. Recrystallisation from a mixture of ethyl acetate and hexane gave analytically pure material of melting point 118-127°C.

15 The 2'-deoxy-5-ethyl-5'-O-triphenylmethyluridine used as the starting material was prepared as follows:

20 A solution of 15.7g of 2'-deoxy-5-ethyluridine and 20.4g of chlorotriphenylmethane in 290ml of dry pyridine was stirred under nitrogen and heated at 100°C for 30 min. The mixture was cooled and poured into 3l of ice/water and extracted with three 500ml portions of ethyl acetate. The combined extracts were washed with 1.5l of water then dried and evaporated. The residue was taken up in 30ml of acetone and 210ml of hot toluene added. The acetone was removed by boiling on a hot water bath. After cooling at -20°C the precipitate was filtered off and washed with diethyl ether to give 19.5g of 2'-deoxy-5-ethyl-5'-O-triphenylmethyluridine of melting point 168-172°C.

25

Example 230

By the procedure of Griffin and Todd, J. Chem. Soc., 1958, 1391 was prepared 5'-benzyloxy-2',3'-dideoxy-5-methyluridine of melting point 140°C (decomposition).

30

Example 231

By the procedure of C. K. Chu et al, J. Med. Chem., 1989, 32, 612 was prepared 2',3'-dideoxy-5-ethyl-3'-iodouridine of melting point 161.5-163.5°C.

5

Example 232

By the procedure of C. K. Chu et al, J. Med. Chem., 1989, 32, 612 was prepared 3'-azido-2',3'-dideoxy-5-ethyluridine of melting point 116-118°C.

Example 233

10 A solution of 2.0 g of 1-(5-O-acetyl-3-azido-2,3-dideoxy-1- β -D-ribofuranosyl)-5-methyl-4-(1,2,4-triazol-1-yl)pyrimidin-2(1H)-one in 23ml of dioxane was treated with 3.5ml of concentrated (32%) aqueous ammonia solution and the mixture stirred at room temperature for 6 hours. The solution was evaporated and the residue dissolved in 36ml of a saturated solution of ammonia in methanol which was stirred at room temperature for 3 days. The residue was extracted several times with boiling ethyl acetate. The combined ethyl acetate extracts were filtered and evaporated. The residue was dissolved in ethanol and the solution concentrated to low volume then diluted with ether. The gum which separated crystallised and the solid was filtered to give 0.47g of 3'-azido-2',3'-dideoxy-5-methylcytidine (T.S. Lin et al, J.Med.Chem., 1983, 26, 1691) of melting point 85-88°C.

15

20

The 1-(5-O-acetyl-3-azido-2,3-dideoxy-1- β -D-ribofuranosyl)-5-methyl-4-(1,2,4-triazol-1-yl)pyrimidin-2(1H)-one used as the starting material was prepared as follows:

- 25 a) A solution of 1.34g of 3'-azido3'-deoxythymidine in 13.5ml of anhydrous pyridine was treated with 0.76ml of acetic anhydride and the mixture stirred at room temperature overnight. 2.5ml of methanol was added and the solution stirred for 30min then evaporated to dryness. The residue was taken up in 125ml of dichloromethane and the solution washed with 50ml of 1m hydrochloric acid, 25ml of saturated sodium hydrogen carbonate solution and 25ml of water then dried over anhydrous sodium sulphate, filtered and
- 30

evaporated to give 1.46g of 5'-O-acetyl-3'-azido-3'-deoxythymidine as a colourless gum which was used without further purification.

- 5 b) A suspension of 1.68g of 1,2,4-triazole in 28ml of anhydrous acetonitrile was stirred and heated to 50°C to give a clear solution. This was removed from the heating bath and stirred while 0.97ml of phosphorus oxychloride was added dropwise during 5 min so that the temperature of the reaction mixture was maintained at 50-52°C. A crystalline white precipitate separated. The mixture was stirred at room temperature for 15 min then cooled to 5°C in ice while 6.42ml of anhydrous triethylamine was added dropwise at 5-10°C during 3
10 min. The mixture was stirred for a further 15 min at room temperature then a solution of 1.68g of crude 5'-O-acetyl-3'-azido-3'-deoxythymidine in 17ml of anhydrous acetonitrile was added over 3 min. The mixture was stirred at room temperature overnight then treated with 4.34ml of triethylamine and 1.08ml of water. The mixture was stirred for 10min then evaporated to dryness and the residue taken up in 125ml of dichloromethane. The solution was washed with saturated sodium hydrogen carbonate solution then evaporated to a yield 2.0g
15 of 1-(5-O-acetyl-3-azido-2,3-dideoxy-1-β-D-ribofuranosyl)-5-methyl-4-(1-(1,2,4-triazol-1-yl)pyrimidin-2(1H)-one as a crystalline solid which was used without further purification.

20

Example 234

By the procedure of G. Gosselin, et al, Patent No. WO 0025799 was prepared 1-(3-deoxy-β-L-threo-pentofuranosyl)-5-fluorocytosine.

25

Example 235

By the procedure of R. Saladino et al, Tetrahedron, 1996, 52, 6759 was prepared 4-methylamino-1-(β-D-ribofuranosyl)pyrimidin-2(1H)-one; mass spectrum (ESI) m/z 258 [M+H]⁺.

Example 236

By the procedure of T. Kulikowski and D. Shugar, Acta. Biochim. Pol., 1979, 26, 145 was prepared 5-fluoro-4-methylamino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one; mass spectrum (ESI) m/z 276[M+H]⁺.

5

Example 237

A solution containing 1.5g of cytidine and 0.86g of 2,5-dimethoxytetrahydrofuran in 10ml glacial acetic acid was heated under nitrogen at 110°C for 1 hour. The solvents were evaporated under low vacuum to give a lilac solid, which was purified by flash chromatography on silica-gel using methanol/dichloromethane (1:19) for the elution to give 90mg of 4-(1-pyrrolyl)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one as a white solid; mass spectrum (ESI) m/z 294 [M+H]⁺.

15

Example 238

A solution of 0.3g of 1-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)-4(3H)-oximinopyrimidin-2(1H)-one in 5ml of anhydrous methanol was treated with 0.2ml of a 1M solution of sodium methoxide in methanol and stirred at room temperature for 24 hours. The mixture was evaporated to dryness and the residue purified by flash chromatography on silica gel using methanol/dichloromethane 1:9 for the elution to give 79mg of 4(3H)-oximino-1-(β -L-ribofuranosyl)pyrimidin-2(1H)-one as a white solid of melting point 138-139°C; mass spectrum m/z 260[M+H]⁺.

The 1-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)-4(3H)-oximinopyrimidin-2(1H)-one used as the starting material was prepared as follows:

a) A mixture of 1.0g of uracil and 1.5g of 1-O-acetyl-2,3,5-tri-O-benzoyl-L-ribose in 50ml of anhydrous acetonitrile was treated with 2.21ml of N,O-bis(trimethylsilyl)acetamide and heated at 76°C under nitrogen until a solution was obtained. To the solution was added 0.98g of trimethylsilyl trifluoromethane sulphonate and heating at 70°C then continued overnight. The mixture was cooled, diluted with 500ml of dichloromethane and washed

three times with 50ml of saturated sodium hydrogen carbonate solution. The dichloromethane solution was washed with brine, dried over anhydrous sodium sulphate, filtered and evaporated to give 1.61g of 2',3',5'-tri-O-benzoyl-L-uridine as a white solid; mass spectrum m/z 557 [M+]⁺.

- 5 b) A solution of 1.80g of 1,2,4-triazole was prepared in 25ml of anhydrous acetonitrile by warming. The solution was stirred at room temperature under nitrogen while 0.86g of phosphorus oxychloride was added. A white suspension was obtained which was cooled to 5°C in ice and treated with 3.46ml of triethylamine during 4 min followed dropwise by a solution of 1.61g of 2',3',5'-tri-O-benzoyl-L-uridine in 25ml of anhydrous acetonitrile during 2 min. The mixture was stirred at room temperature for 2.5 hours then treated with a further 2.41ml of triethylamine followed by 0.63ml of water and stirred for 10 min. The mixture was diluted with 150ml of dichloromethane and washed with a 10% solution of sodium hydrogen carbonate and brine. The dichloromethane solution was dried over anhydrous sodium sulphate, filtered and evaporated to give 1.6g of a yellow powder. This was purified by flash chromatography on silica gel using ethyl acetate/isohexane (1:9) for the elution to give 1.12 of 1-(2,3,5-tri-O-benzoyl-β-L-ribofuranosyl)-4-(1,2,4-triazol-1-yl)pyrimidin-2(1H)-one as a white solid of melting point 83-84°C; mass spectrum m/z 608 [M+H]⁺.
- 10 c) A suspension of 0.43g of hydroxylamine hydrochloride in 15ml of anhydrous methanol was treated with 4.96ml of a 1M solution of sodium methoxide in methanol. After stirring for 10 min a solution of 0.75g of 1-(2,3,5-tri-O-benzoyl-β-L-ribofuranosyl)-4-(1,2,4-triazol-1-yl)pyrimidin-2(1H)-one in a mixture of 20ml of methanol and 20ml of tetrahydrofuran was added and the mixture stirred at room temperature overnight. The mixture was evaporated and the residue purified by flash chromatography on silica gel using methanol/dichloromethane 1:24 for the elution to give 0.605g of 1-(2,3,5-tri-O-benzoyl-β-L-ribofuranosyl)-4(3H)-oximinopyrimidin-2(1H)-one as a white solid; mass spectrum m/z 572[M+H]⁺.
- 15
- 20
- 25
- 30

Example 239

In an analogous manner to that described in example 238 was prepared 4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one (I.Wempen et al, J.Med.Chem.,1968, 11, 144); mass spectrum (ESI) m/z 260[M+H]⁺.

5

Example 240

In an analogous manner to that described in example 238 was prepared 4-oximino-1-(β -D-arabinofuranosyl)pyrimidin-2(1H)-one, (I.Wempen et al, J.Med.Chem.,1968, 11, 144).

10

Example 241

In an analogous manner to that described in example 238 was prepared 5-fluoro-4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one; mass spectrum m/z 319 [M+H]⁺.

15

Example 242

By the procedure of S. L. Anliker et al, J. Pharm. Sci., 1994, 83, 716 was prepared 1-(2-deoxy-2,2-difluoro- α -D-erythropentofuranosyl)uracil.

20

Example 243

By the procedure of S. L. Anliker et al, J. Pharm. Sci., 1994, 83, 716 was prepared 1-(2-deoxy-2,2-difluoro- β -D-erythropentofuranosyl)cytosine.

25

Example 244

By the procedure of E Moyroud and P Strazewski, Tetrahedron, 1999, 55, 1277 was prepared L-cytidine or according the following experimental method:

5 A solution of 0.40g of 1-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)-4-(1H-1,2,4-triazol-1-yl)-1H-pyrimidin-2-one in 10ml of 1,4-dioxane was treated with 0.5ml of 35% aqueous ammonia solution and stirred at room temperature for 12 hours. The reaction mixture was concentrated under reduced pressure to leave a white solid which was purified by flash chromatography on silica gel using dichloromethane/methanol (1:24 then 1:9) to give 0.22g of 2',3',5'-tri-O-benzoyl-L-cytidine. This was dissolved in 2ml of anhydrous methanol and treated with 100 μ l of 1M sodium methoxide solution. The reaction mixture was stirred for 16 hours then evaporated and the residue purified by flash chromatography on silica gel using dichloromethane/methanol(9:1 then 3:2) for the elution to give 80mg of L-cytidine as a white solid; mass spectrum(ESI) m/z 301 [M+H+MeCN]⁺.

10

The 1-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)-4-(1H-1,2,4-triazol-1-yl)-1H-pyrimidin-2-one used as the starting material was prepared as follows:

- 15 a) A mixture containing 1.0g of uracil and 1.5g of 1-O-acetyl-2,3,5-tri-O-benzoyl-L-ribose in 50ml of anhydrous acetonitrile was treated with 1.82g of N,O-bis-trimethylsilylacetamide and heated at 76°C under nitrogen until a clear solution was obtained. 0.98g of trimethylsilyl trifluoromethanesulphonate was then added in one portion and heating at 70°C continued for 16 hours. The mixture was cooled and diluted with 500ml of dichloromethane. The solution was washed three times with 50ml of saturated sodium hydrogen carbonate solution and brine then dried over anhydrous sodium sulphate, filtered and evaporated to give 1.61g of 2',3',5'-tri-O-benzoyl-L-uridine as a white solid which was used without further purification.
- 20 b) To a solution of 1.80g of 1,2,4-triazole in 25ml of anhydrous acetonitrile at room temperature under nitrogen was added 0.86g of phosphorus oxychloride. The mixture was cooled in a bath of ice and stirred for 15 min then treated with 2.53g (3.46ml) of triethylamine during 4 min. The ice bath was removed and a solution of 1.61g of 2',3',5'-tri-O-benzoyl-L-uridine in 25ml of anhydrous acetonitrile added dropwise during 2 min. The reaction mixture was stirred at room temperature under nitrogen for 2.5 hours then a further 2.41ml of triethylamine added followed by 0.63ml of water. After stirring for 10 min the reaction mixture was diluted with 150ml of dichloromethane and washed with a 10% aqueous solution of sodium hydrogen carbonate. The dichloromethane solution was washed with brine then dried over anhydrous sodium sulphate. Evaporation gave 1.61g of solid which was purified by flash chromatography on
- 25
- 30
- 35

silica gel using ethyl acetate/isohexane (1:9) for the elution to give 1.12g of 1-(2,3,5-tri-O-benzoyl- β -L-ribofuranosyl)-4-(1H-1,2,4-triazol-1-yl)-1H-pyrimidin-2-one as a white solid; mass spectrum (ESI) m/z 608 [M+H]⁺.

5

Example 245

A solution of 55mg of 1-[1(R)-2,2-difluoro-3(R)-acetoxy-4(R)-(acetoxymethyl)-cyclopentyl]-4-(1H-1,2,4-triazol-1-yl)-1H-pyrimidin-2-one in 35% aqueous ammonia was stirred at room temperature for 14 hours. The reaction mixture was concentrated under reduced pressure and the residue purified by chromatography on silica gel using dichloromethane/methanol (5:1) for the elution to give 35mg of 1-[1(R)-2,2-difluoro-3(R)-hydroxy-4(R)-(hydroxymethyl)cyclopentyl]-4-amino-1H-pyrimidin-2-one as colourless crystals; mass spectrum (ESI) m/z 262 [M+H]⁺; ¹H NMR (270MHz, DMSO-d₆) 1.71 (1H, m), 1.90 (1H, m), 1.99 (1H, m), 3.45 (1H, m), 3.55 (1H, m), 3.80 (1H, m), 4.73 (1H, t), 5.22 (1H, m), 5.68 (1H, d), 5.71 (1H, d), 7.15 (1H, br.s), 7.18 (1H, br.s), 7.56 (1H, d).

The 1-[1(R)-2,2-difluoro-3(R)-acetoxy-4(R)-(acetoxymethyl)-cyclopentyl]-4-(1H-1,2,4-triazol-1-yl)-1H-pyrimidin-2-one used as the starting material was prepared as follows:

- a) A solution of 28g of (3aS,4R,7S,7aR)-tetrahydro-2,2-dimethyl-4,7-methano-1,3-dioxolo[4,5-c]pyridin-6(3aH)-one, in 300ml of 10% methanolic hydrogen chloride was stirred at ambient temperature for 3 days. The reaction mixture was concentrated under reduced pressure to ca 100ml and cooled in a refrigerator. The white precipitate was collected and washed with methanol to give a first crop of 25.44g of (1S,2R,3S,4R)-4-amino-2,3-dihydroxy-cyclopentanecarboxylic acid methyl ester hydrochloride. The combined mother liquor and washings were concentrated and recrystallised from methanol to give 4.30g of a second crop; ¹H NMR (270MHz, DMSO-d₆) 1.68 (1H, dddd), 2.22 (1H, dddd), 3.2-3.35 (1H, br.m), 3.62 (3H, s), 3.80-3.90 (1H, br.m), 4.00-4.10 (1H, br.m), 5.20 (1H, br.s), 5.30 (1H, br.s), 8.39 (3H, br.s).
- b) To a solution of 28.6g of (1S,2R,3S,4R)-4-amino-2,3-dihydroxy-cyclopentanecarboxylic acid methyl ester hydrochloride and 35.36g of di-t-butyl dicarbonate in 400ml of dioxane was added 27.2g of sodium hydrogen

carbonate dissolved in a minimum volume of water and the reaction mixture was stirred at ambient temperature for 36 hours. The reaction mixture was filtered and the filter washed thoroughly with 300ml of acetone. The filtrate and washings were concentrated under reduced pressure to ca 100ml and the residue partitioned between 300ml of ethyl acetate and 100ml of water. The water layer was extracted further with 300ml of ethyl acetate. The combined organic extracts were dried over anhydrous sodium sulphate, filtered and concentrated under reduced pressure. The residue was recrystallised from 200ml of diethyl ether to give 34.9g of (1S,2R,3S,4R)-4-t-butoxycarbonylamino-2,3-dihydroxy-cyclopentanecarboxylic acid methyl ester as colourless crystals; H¹ NMR (270MHz, CDCl₃) 1.45 (9H, s), 1.60-1.75 (1H, m), 2.35-2.45 (1H, m), 2.93 (1H, ddd), 3.10 (1H, br.s), 3.71 (3H, s), 3.80-3.95 (2H, m), 4.28 (1H, m), 4.65 (1H, br.s), 4.88 (1H, br.s).

c) To a solution of 33.77g of (1S,2R,3S,4R)-4-t-butoxycarbonylamino-2,3-dihydroxy-cyclopentanecarboxylic acid methyl ester in 300ml of anhydrous tetrahydrofuran was added dropwise a solution of 4.0g of lithium borohydride in 100ml of anhydrous tetrahydrofuran and the reaction mixture stirred for 2 hours at ambient temperature. The excess lithium borohydride was decomposed by addition of 10ml of water and stirring for a short time. The reaction mixture was dried over anhydrous sodium sulphate, filtered and the filter washed thoroughly with tetrahydrofuran. The combined filtrate and washings were concentrated under reduced pressure and dried under vacuum to give crude (1R,2S,3R,5R)-3-t-butoxycarbonylamino-5-hydroxymethyl-cyclopentan-1,2-diol which was redissolved in 100ml of dioxane and treated dropwise with 300ml of a 4M solution of hydrogen chloride in dioxane. The reaction mixture was stirred at ambient temperature for 14 hours. The solvent and volatile materials were removed by purging with nitrogen gas and then evaporation under reduced pressure. The residue was rinsed twice with 100ml of n-hexane then dried under vacuum to give crude (1R,2S,3R,5R)-3-amino-5-hydroxymethyl-cyclopentan-1,2-diol hydrochloride. A solution of this and 22.8g of 2,4-dinitro-fluorobenzene in 100ml of absolute N,N-dimethyl formamide was treated with sodium hydrogen carbonate and the suspension stirred at ambient temperature for 5 hours. The reaction mixture was filtered and the filter washed thoroughly with methanol. The combined filtrate and washings were concentrated under reduced pressure and the residue purified by chromatography on silica gel using dichloromethane /methanol(9:1 to 4:1)

for the elution to give 30.15g of (1R,2S,3R,5R)-3-[(2,4-dinitrophenyl)amino]-5-hydroxymethyl-cyclopentan-1,2-diol as an amorphous yellow solid; ¹H NMR (270MHz, DMSO-d₆) 1.32 (1H, ddd), 1.95-2.05 (1H, m), 2.35 (1H, ddd), 3.44 (2H, s), 3.70-3.85 (2H, m), 3.99 (1H, ddd), 4.62 (1H, br.t), 4.78 (1H, br.d), 5.03 (1H, br.d), 7.33 (1H, d), 8.27 (1H, dd), 8.67 (1H, d), 8.86 (1H, d).

- 5
- d) To a solution of 30.15g of (1R,2S,3R,5R)-3-[(2,4-dinitrophenyl)amino]-5-hydroxymethyl-cyclopentan-1,2-diol and 19.69g of imidazole in 150ml of dry N,N-dimethylformamide was added in portions tetra-isopropyl dichlorosiloxane. The reaction mixture was stirred at room temperature under argon for 14 hours then poured into 500ml of water and extracted twice with 400ml of ethyl acetate. The combined organic extracts were washed twice with 300ml of brine, dried over anhydrous sodium sulphate, filtered and evaporated to give yellow sticky crystals which were recrystallised from n-hexane to give 43.23g of 2[(2,4-dinitrophenyl)amino-5,5,7,7-tetraisopropyl-hexahydro-4,6,8-trioxa-5,7-disila-cyclopentacyclooctene-3-ol in two crops.
- 10
- e) To a solution of 2.0g of 2[(2,4-dinitrophenyl)amino-5,5,7,7-tetraisopropyl-hexahydro-4,6,8-trioxa-5,7-disila-cyclopentacyclooctene-3-ol in 15ml of dry acetonitrile was added 4.0g of 1,1,1-triacetoxy-1,1-dihydro-1,2-benziodoxol-3(1H)-one and the suspension stirred at 40°C under argon for 14 hours. The reaction mixture was diluted with 40ml of saturated sodium hydrogen carbonate solution and extracted twice with 50ml of dichloromethane. The combined organic extracts were washed successively with 40ml of saturated sodium hydrogen carbonate solution and 40ml of brine then dried over anhydrous sodium sulphate, filtered and evaporated. The yellow amorphous residue which was purified by chromatography on silica gel using n-hexane/ethyl acetate (4:1) for the elution to give 1.50g of 2-[(2,4-dinitrophenyl)amino)-5,5,7,7-tetraisopropyl-hexahydro-4,6,8-trioxa-5,7-disila-cyclopentacyclooctene-3-one; ¹H NMR (270MHz, CDCl₃) 1.02-1.15 (28H, m), 1.55-1.62 (1H, br.m), 2.16-2.28 (1H, m), 2.54-2.66 (1H, m), 3.93 (1H, dd), 4.14 (1H, dd), 4.20 (1H, m), 4.30 (1H, d), 7.17 (1H, d), 8.28 (1H, dd), 8.63 (1H, br.d), 9.14 (1H, d).
- 15
- f) To an ice-cooled solution of 6.24ml of diethylamino sulphur trifluoride complex in 24ml of dry dichloromethane was added dropwise over 10 min a solution of 2.0g of 2-[(2,4-dinitrophenyl)amino)-5,5,7,7-tetraisopropyl-hexahydro-4,6,8-trioxa-5,7-disila-cyclopentacyclooctene-3-one in 24ml of dry
- 20
- 25
- 30
- 35

- dichloromethane. The mixture was stirred at 0°C under argon for 4 hours then poured into 100ml of sodium hydrogen carbonate solution and extracted three times with 100ml of dichloromethane. The combined extracts were washed successively with three portions of 200ml of sodium bicarbonate solution and twice with 100ml of brine then dried over anhydrous sodium sulphate, filtered and evaporated . The dark yellow amorphous residue was purified by chromatography on silica gel using n-hexane/dichloromethane (1:1) for the elution to give 0.59g of (3,3-difluoro-5,5,7,7-tetraisopropyl-hexahydro-4,6,8-trioxa-5,7-disilacyclopentacyclooctene-2-yl)(2,4-dinitrophenyl)amine; ¹H NMR (270MHz, CDCl₃) 1.02-1.15 (28H, m), 1.60-1.72 (1H, br.m), 2.02-2.16 (1H, m), 2.36-2.48 (1H, m), 3.80 (1H, dt), 4.05 (1H, dd), 4.10-4.20 (2H, m), 7.05 (1H, d), 8.28 (1H, dd), 8.50 (1H, br.d), 9.14 (1H, d).
- g) To an ice-cooled solution of 0.677g of (3,3-difluoro-5,5,7,7-tetraisopropyl-hexahydro-4,6,8-trioxa-5,7-disilacyclopentacyclooctene-2-yl)-(2,4-dinitrophenyl)amine in 15ml of tetrahydrofuran was added 2.5ml of a 1M solution of tetrabutylammonium fluoride in tetrahydrofuran and the reaction mixture was stirred at 0°C under an atmosphere of argon for 3 hours. The reaction mixture was concentrated under reduced pressure and the residue was partitioned between 40ml of ethyl acetate and 50ml of water. The water layer was extracted further with three portions of 40ml of ethyl acetate. The combined extracts were washed with 30ml of brine , dried over anhydrous sodium sulphate, filtered and evaporated. The residue was purified by chromatography on silica gel using dichloromethane/ methanol (20:1) for the elution to give 0.364g of (1R,3R,5R)-3-[(2,4-dinitrophenyl)amino]-2,2-difluoro-5-(hydroxymethyl)cyclopentanol as a pale yellow solid; ¹H NMR (270MHz, CDCl₃) 1.70 (1H, m), 2.22 (1H, m), 2.54 (1H, m), 3.70-3.90 (3H, m), 4.18 (1H, m), 4.40 (1H, m), 4.66 (1H, d), 7.21 (1H, d), 8.31 (1H, dd), 8.78 (1H, br.d), 9.11 (1H, d).
- h) To a solution of 0.36g of (1R,3R,5R)-3-[(2,4-dinitrophenyl)amino]-2,2-difluoro-5-(hydroxymethyl)cyclopentanol in 20ml of 75% aqueous acetone was treated with 1.0g of Dowex-1 ion-exchange resin , which had been thoroughly washed successively with 1M sodium hydroxide solution, distilled water and methanol prior to use. The reaction mixture was stirred at ambient temperature for 24 hours. The resin was filtered off and thoroughly washed with approximately 100ml of 75% aqueous acetone. The combined filtrate was concentrated under reduced pressure to remove acetone and the resulting

aqueous solution acidified with 2ml 1M hydrochloric acid. The aqueous solution was washed twice with 20ml of ethyl acetate then lyophilised to give 0.134g of (1R,3R,5R)-3-amino-2,2-difluoro-5-(hydroxymethyl)-cyclopentanol hydrochloride as a colourless powder.

- 5 i) To a solution of 0.127g of (1R,3R,5R)-3-amino-2,2-difluoro-5-(hydroxymethyl)-cyclopentanol hydrochloride in 2ml of anhydrous N,N-dimethylformamide were added freshly desiccated 4°A molecular sieves. The mixture was stirred at 30°C for 30 minutes then treated with 2.5ml of a 0.427M solution of 3-ethoxy-2-propenoyl isocyanate. The mixture was stirred at 30°C for 30 minutes and then at room temperature for 14 hours. The reaction mixture was concentrated under reduced pressure and the residue purified by chromatography on silica gel using dichloromethane/methanol(9:1 then 5:1)) for the elution to give 0.157g of 1-[1(R)-2,2-difluoro-3(R)-hydroxy-4(R)-(hydroxymethyl)cyclopentyl]-3(3-ethoxy-E-2-propenoyl)urea as a colourless solid; ¹H NMR (270MHz, DMSO-d₆) 1.23 (3H, t), 1.78-1.91 (1H, m), 2.04-2.15 (1H, m), 3.38-3.45 (2H, m), 3.60-3.75 (1H, m), 3.96 (2H,q), 4.22-4.44 (1H, m), 4.72 (1H, t), 5.51 (1H, d), 5.63 (1H, d), 7.59 (1H, d), 8.80 (1H, s). 10.21 (1H, s).
- 10 j) A solution of 0.15g of 1-[1(R)-2,2-difluoro-3(R)-hydroxy-4(R)-(hydroxymethyl)cyclopentyl]-3(3-ethoxy-E-2-propenoyl)urea in 4ml of 5% aqueous sulphuric acid was boiled under reflux for 3 hours. The reaction mixture was neutralised by addition of sodium hydroxide solution then concentrated under reduced pressure. The residue was suspended in 35ml of absolute ethanol and filtered. The material on the filter was washed three times with 35ml of absolute ethanol and the combined filtrate concentrated under reduced pressure to give 0.215g of crude 1-[1(R)-2,2-difluoro-3(R)-hydroxy-4(R)-(hydroxymethyl)cyclopentyl]-1H-pyrimidine-2,4-dione as a colourless powder which was used without further purification.
- 15 k) To a solution of 0.215g of crude 1-[1(R)-2,2-difluoro-3(R)-hydroxy-4(R)-(hydroxymethyl)cyclopentyl]-1H-pyrimidine-2,4-dione in 3ml of acetic anhydride was added 5mg of 4-dimethylaminopyridine and the reaction mixture stirred at room temperature for 14 hours. The mixture was concentrated under reduced pressure and the residue partitioned between 30ml of ethyl acetate and sodium hydrogen carbonate solution. The aqueous layer was extracted twice more with 30ml of ethyl acetate. Combined extracts were washed with 30ml of brine, dried over anhydrous sodium sulphate, filtered and
- 20
- 25
- 30
- 35

evaporated. The residue was triturated with t-butyl methyl ether to give 0.148g of 1-[1(R)-2,2-difluoro-3(R)-acetoxy-4(R)-(acetoxymethyl)cyclopentyl]-1H-pyrimidine-2,4-dione, which was used without further purification.

- l) To a solution of 0.128g of 1-[1(R)-2,2-difluoro-3(R)-acetoxy-4(R)-(acetoxymethyl)cyclopentyl]-1H-pyrimidine-2,4-dione and 0.128g of 1,2,4-1H-triazole in dry pyridine was added dropwise 180 μ l of 4-chlorophenyl dichlorophosphate and the reaction mixture stirred at room temperature for 14 hours. The reaction mixture was concentrated under reduced pressure and the residue partitioned between 30ml of ethyl acetate and sodium hydrogen carbonate solution. The aqueous layer was extracted twice more with 30ml of ethyl acetate and the combined extracts washed with 30ml of brine, then dried over anhydrous sodium sulphate, filtered and evaporated. The residue was purified by chromatography using ethyl acetate for the elution to give 0.112g of 1-[1(R)-2,2-difluoro-3(R)-acetoxy-4(R)-(acetoxymethyl)cyclopentyl]-4-(1H-1,2,4-triazol-1yl)-1H-pyrimidin-2-one; 1 H NMR (270MHz, CDCl₃) 1.86 (1H, m), 2.11 (3H, s), 2.18 (3H, s), 2.30-2.50 (1H, m), 2.50-2.70 (1H, m), 4.15-4.25 (2H, m), 5.19 (1H, ddd), 5.55-5.75 (1H, m), 7.12(1H, d), 7.93 (1H, d), 8.15 (1H, s), 9.29 (1H, s).

20

Example 246

Starting with 1(R)-amino-2(S),3(R)-diacetoxy-4(R)-acetoxymethylcyclopentane in manner analogous to that described by Y. F. Shealy and C. A. O'Dell, J. Heterocyclic Chem., 1980, 17, 353 was prepared 4-amino-1(R)-(2(S),3(R)-dihydroxy-4(R)-hydroxymethyl-cyclopentyl)-1H-pyrimidin-2-one; mass spectrum(ESI) m/z 242 [M+H]⁺.

25

Example 247

The compound may be prepared according to G. Gosselin et al, J. Med. Chem. 1987, 30960, 982. A solution of 0.283g of 1-(3-O-acetyl-2,5-bis-O-t-butyldimethylsilyl- β -D-xylofuranosyl)cytosine and 0.245g of ammonium fluoride in 5 ml of anhydrous methanol was stirred and heated at 50-60°C under nitrogen for 24 hours. The solution was evaporated and the white solid residue purified by

flash chromatography on silica gel using methanol/dichloromethane (1:19 to 2:3) for the elution to give 50mg of 1-(β -D-xylofuranosyl)cytosine; mass spectrum(ESI) m/z 244 [M+H]⁺.

The 3-O-acetyl-2,5-bis-O-t-butyldimethylsilyl-1-(β -D-xylofuranosyl)cytosine used as the starting material was prepared as follows:

- a) A solution of 0.5g of 1-(2,5-bis-O-t-butyldimethylsilyl- β -D-xylofuranosyl)uracil (prepared according to F. Hansske, D. Madej and M. J. Robins, Tet., 1984, 40, 125) in 5ml of anhydrous pyridine was treated with 120 μ l of acetic anhydride and stirred at room temperature for 30 hours. A further 120 μ l of acetic anhydride was added and stirring continued for a further 3 days. The reaction mixture was treated with 0.2 ml of water and then evaporated. The pale yellow oily residue was taken up in 70 ml of dichloromethane and the solution washed with three 10 ml portions of 1M hydrochloric acid then dried over anhydrous sodium sulphate , filtered and evaporated to give 0.53g of 1-(3-O-acetyl-2,5-bis-O-t-butyldimethylsilyl- β -D-xylofuranosyl)uracil as a pale yellow oil which was used without further purification.
- b) A solution of 0.629g of 1,2,4-triazole in 15 ml of anhydrous acetonitrile was treated with 182 μ l of phosphorus oxychloride. A white suspension formed which was cooled in ice for 15 min then treated with 1.21 ml of triethylamine. The ice bath was removed while a solution of 0.52 g of 1-(3-O-acetyl-2,5-bis-O-t-butyldimethylsilyl- β -D-xylofuranosyl)uracil in 10 ml of dry acetonitrile was added dropwise over 3 minutes. The reaction mixture was stirred at room temperature under nitrogen overnight then diluted with dichloromethane and washed with saturated sodium hydrogen carbonate solution. The dichloromethane solution was dried over anhydrous sodium sulphate, filtered and evaporated. The residue was purified by flash chromatography on silica gel using ethyl acetate/isohexane (1:9 to 3:5) for the elution to give 0.286g of 1-(3-O-acetyl-2,5-bis-O-tert-butyldimethylsilyl- β -D-ribofuranosyl)-4-(1-triazolyl)pyrimidin-2(1H)-one; mass spectrum(ESI) m/z 566 [M+H]⁺.
- c) A solution of 0.28g of 1-(3-O-acetyl-2,5-bis-O-t-butyldimethylsilyl- β -D-ribofuranosyl)-4-(1-triazolyl)pyrimidine-2(1H)-one in 10 ml of 1,4-dioxane was treated with 0.5 ml of concentrated aqueous ammonia solution and stirred at room temperature for 12 hours then evaporated to yield 0.25g of 1-(3-O-

acetyl-2,5-bis-O-t-butyldimethylsilyl- β -D-xylofuranosyl)cytosine as a white solid; mass spectrum(ESI) m/z 514 [M+H]⁺.

Example 248

5 The compound may be prepared according to H. Hayakawa et al, Chem. Pharm.Bull., 1990, 38(5), 1136. A mixture of 0.3g of 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)uracil (prepared according to H. Hayakawa et al, Chem. Pharm. Bull., 1990, 38, 1136) and 80% acetic acid was stirred and heated at 100° C for 5 hours then evaporated to dryness. The residue was
10 redissolved in 10 ml of distilled water and the solution washed with three 5ml portions of diethyl ether. The aqueous solution was evaporated to dryness and the residue purified by flash chromatography on silica gel using methanol/dichloromethane (1:19 to 1:12) for the elution to give 53 mg of 1-(3-deoxy-3-fluoro- β -D-xylofuranosyl)uracil; mass spectrum(CI) m/z 246 [M+H]⁺.

15

Example 249

The compound may be prepared according to J. A. Wright, D. P. Wilson and J. J. Fox, J. Med. Chem. 1970, 13(2), 269. A solution of 0.2g of 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)cytosine in 5 ml of dry methanol was stirred with 1.2g of Amberlyst 15 ion exchange resin for 5 hours. The resin was filtered off and washed with methanol then suspended in 10ml of methanol/1M ammonia solution(1:1) and stirred for 30 min. The mixture was filtered and the resin washed thoroughly with methanol. The filtrate was evaporated to a glass which was purified by flash chromatography on silica gel using methanol/dichloromethane (1:4) for the elution to give 13 mg of 1-(3-deoxy-3-fluoro- β -D-xylofuranosyl)cytosine; mass spectrum(ESI) m/z 246 [M+H]⁺.

20

25

The 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)cytosine used as the starting material was prepared as follows:

30

- a) A solution of 1.71g of 1,2,4-triazole in 20ml of anhydrous acetonitrile was stirred under nitrogen and treated with 0.47 ml of phosphorus oxychloride to give a milky suspension which was cooled to <5°C for 15 min then treated with 3.2 ml of triethylamine. After allowing to warm to room temperature a

suspension of 2.0g of 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)uracil (prepared according to H. Hayakawa et al, Chem. Pharm. Bull., 1990, 38, 1136) in 15 ml of acetonitrile was added and the mixture stirred at room temperature for 24 hours. The mixture was diluted with dichloromethane and washed with saturated sodium hydrogen carbonate solution. The dichloromethane solution was dried over anhydrous sodium sulphate, filtered and evaporated. The residue was purified by flash chromatography on silica gel using ethyl acetate/isohexane (1:1) for the elution to give 0.5g of 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)-4-(1,2,4-triazolyl)pyrimidin-2(1H)-one as a white solid; mass spectrum (ESI) m/z 782 [M+H]⁺.

10 b) A solution of 0.5g of 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)-4-(1,2,4-triazolyl)pyrimidin-2(1H)-one in 10 ml of 1,4-dioxane was treated with 1 ml of concentrated ammonia solution and stirred at room temperature for 16 hours. The solution was evaporated to dryness and the residue purified by flash chromatography on silica gel using ethyl acetate/iso hexane (1:1) for the elution to give 0.23g of 1-(3-deoxy-3-fluoro-2,5-bis-O-triphenylmethyl- β -D-xylofuranosyl)cytosine; mass spectrum (CI) m/z 731 [M+H]⁺.

20

Example 250

The compound may be prepared according to R. Z. Sterzycki, M. M. Mansuri and J. C. Martin, Eur. Pat. Appl. (1990) EP 391411. A solution of 55mg of 4-N-acetyl-1-(3-acetoxymethyl-2,3-di-O-acetyl-3-deoxy- β -D-ribofuranosyl)cytosine in 0.5 ml of anhydrous methanol was treated with 0.05ml of 1M sodium methoxide solution and stirred at room temperature for 5 hours. The solution was neutralised by addition of a few drops of glacial acetic acid and evaporated. The residue was purified by recrystallisation from methanol/ethyl acetate to give 3'-deoxy-3'-hydroxymethylcytidine as a white solid; mass spectrum (ESI) m/z 258[M+H]⁺.

25

30 The 4-N-acetyl-1-(3-acetoxymethyl-2,3-di-O-acetyl-3-deoxy- β -D-ribofuranosyl)cytosine used as the starting material was prepared as follows:

A mixture of 0.3 g of 3-acetoxymethyl-1,2,5-tri-O-acetyl-3-deoxy- β -D-ribofuranose (prepared by the procedure of R. M. Sterzycki et al Eur.Pat.Appl. 391411), 0.457g of N-acetylcytosine and 0.74 ml of bis-trimethylsilylacetamide in 15 ml of anhydrous acetonitrile was heated under reflux for 2.5 hours to give a clear solution. The solution was cooled and treated with 0.28 ml of trimethylsilyl trifluoromethanesulphonate then heated at 50°C for 3 days. The pale yellow solution was diluted with 100 ml of ethyl acetate and washed with 50 ml of 1M hydrochloric acid, 50 ml of saturated sodium hydrogen carbonate then brine. The solution was dried over anhydrous magnesium sulphate, filtered and evaporated.

The residue was purified by flash chromatography on silica gel using methanol/dichloromethane (1:19) for the elution to give 55 mg of 4-N-acetyl-1-(3-acetoxymethyl-2,3-di-O-acetyl-3-deoxy- β -D-ribosyl)cytosine; mass spectrum (ESI) 426[M+H]⁺.

15

Example 251

The compound may be prepared according to R. Z. Sterzycki, M. M. Mansuri and J. C. Martin, Eur. Pat. Appl. (1990) EP 391411. 2'-Deoxy-2'-methoxyuridine is available commercially from ICN Biomedicals Inc., Cat. No. 104991.

20

Example 252

The compound may be prepared according to E. Lescrinier et al, Nucleosides and Nucleotides, 1996, 15, 1863. In a manner analogous to that described in Example 38 starting with 6-chloro-9-(β -D-ribosyl)purine was prepared 6-ethylamino-9-(β -D-ribosyl)purine; mass spectrum(ESI) m/z 296 [M+H]⁺.

25

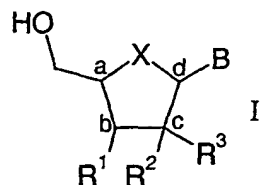
Example 253

The compound may be prepared according to E. Lescrinier et al, Nucleosides and Nucleotides, 1996, 15, 1863. In a manner analogous to that described in Example 38 starting with 6-chloro-9-(β -D-ribosyl)purine was prepared 6-propylamino-9-(β -D-ribosyl)purine; mass spectrum(ESI) m/z 310 [M+H]⁺.

- It will be understood that references herein to treatment extend to prophylaxis as well as to the treatment of existing conditions, and that the treatment of animals includes the treatment of humans as well as other mammals.
- 5 Furthermore, treatment of an Hepatitis C Virus (HCV) infection, as used herein, also includes treatment or prophylaxis of a disease or a condition associated with or mediated by Hepatitis C Virus (HCV) infection, or the clinical symptoms thereof.
- In the present specification "comprise" means "includes or consists of" and "comprising" means "including or consisting of".
- 10 The features disclosed in the foregoing description, or the following claims, or the accompanying drawings, expressed in their specific forms or in terms of a means for performing the disclosed function, or a method or process for attaining the disclosed result, as appropriate, may, separately, or in any combination of such features, be utilised for realising the invention in diverse forms thereof.
- 15

Claims

1. Use of compounds of formula I



5 wherein

R^1 is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy, halogen, cyano, isocyano or azido;

R^2 is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine;

R^3 is hydrogen; or

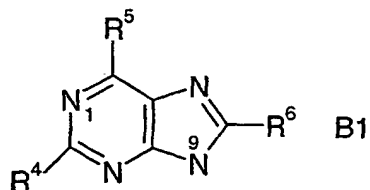
10 R^2 and R^3 together represent $=CH_2$; or

R^2 and R^3 represent fluorine;

X is O, S or CH_2 ;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

15 B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

R^4 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen or SH;

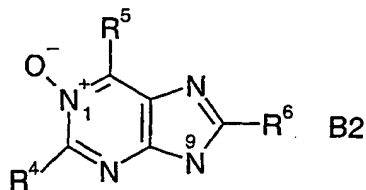
R^5 is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclylamino, halogen, NR^7R^8 , $NHOR^9$, $NHNR^7R^8$ or SH ;

5 R^6 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen, SH or cyano;

R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R^9 is hydrogen, alkyl or aryl; or

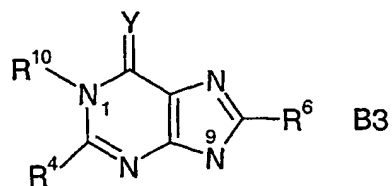
10 B signifies an oxidised purine base B2 which is connected through the 9-nitrogen of formula



wherein

R^4 , R^5 and R^6 are as defined above; or

B signifies a purine base B3 which is connected through the 9-nitrogen of formula



15

wherein

R^4 and R^6 are as defined above;

R^{10} is hydrogen, alkyl or aryl;

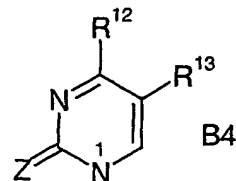
Y is O, S or NR^{11} ;

- 178 -

R¹¹ is hydrogen, hydroxy, alkyl, OR⁹, heterocycl or NR⁷R⁸;

R⁷, R⁸ and R⁹ are as defined above; or

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



5

wherein

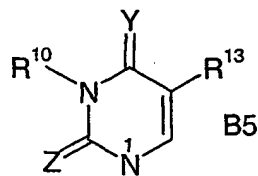
Z is O or S;

R¹² is hydrogen, hydroxy, alkyl, alkoxy, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH;

10 R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen;

R⁷, R⁸ and R⁹ are as defined above; or

B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula



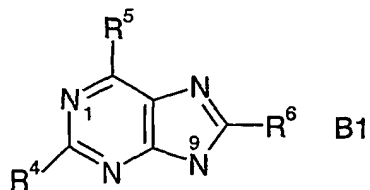
15 wherein

Y, Z, R¹⁰ and R¹³ are as defined above;

for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

20 2. Use of compounds of formula I as claimed in claim 1 wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula

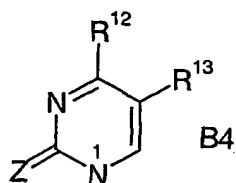


wherein

R^4, R^5, R^6, R^7, R^8 and R^9 are as defined in claim 1;

5 with the proviso that R^4 is not NH_2 and R^5 is not $NH(CH_3)$; or

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula

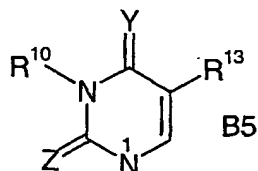


wherein

10 $Z, R^7, R^8, R^9, R^{12}, R^{13}$ are as defined in claim 1;

with the proviso that R^{12} is not hydroxy, alkoxy, $N(CH_3)_2$, $N(H)NH(CH_3)$ or $N(H)NH_2$ and R^{13} is not hydroxyalkyl, chlorine or bromine; or

B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula



15

wherein

Y, Z, R^{10} and R^{13} are as defined in claim 1;

- 180 -

with the proviso that R¹⁰ is not methyl or hydroxyethyl.

3. Use of compounds of formula I as claimed in claims 1 or 2 wherein

R¹ is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy, halogen;

5 R² is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine;

R³ is hydrogen; or

R² and R³ represent fluorine;

X is O;

a, b, c and d denoting asymmetric carbon atoms and forming a D-ribofuranosyl

10 ring.

4. Use of compounds of formula I as claimed in any one of claims 1 to 3
wherein

R¹ is hydroxy;

15 R² is hydroxy;

R³ is hydrogen; or

X is O;

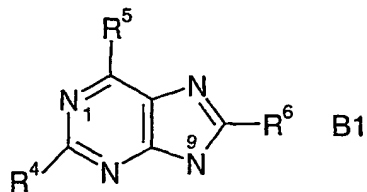
a, b, c and d denoting asymmetric carbon atoms and forming a β-D-ribofuranosyl
ring.

20

5. Use of compounds of formula I as claimed in any one of claims 1, 3 or 4
wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula

- 181 -



wherein

R^4 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen or SH;

5 R^5 is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclylamino, halogen, NR^7R^8 , $NHOR^9$, $NHNR^7R^8$ or SH;

R^6 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen, SH or cyano;

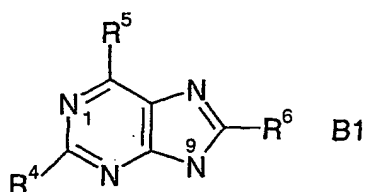
10 R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R^9 is hydrogen, alkyl or aryl.

6. Use of compounds of formula I as claimed in any one of claims 1 or 3 to 5

15 wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

R^4 is hydrogen, chlorine or NH_2 ;

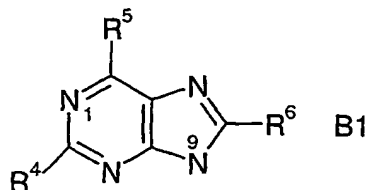
20 R^5 is hydroxy, alkylthio, aryl, heterocyclyl, halogen, NR^7R^8 or SH;

R^6 is hydrogen, halogen, heterocyclyl or NR^7R^8 ;

R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, alkenylalkyl or alkynylalkyl.

- 5 7. Use of compounds of formula I as claimed in any one of claims 1 or 3 to 6
wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

10 R^4 is hydrogen;

R^5 is alkylthio, aryl, heterocyclyl, halogen or NR^7R^8 ;

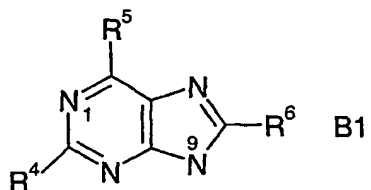
R^6 is hydrogen or halogen;

R^7 and R^8 are independently of each other hydrogen, alkyl, alkenylalkyl or alkynylalkyl.

15

8. Use of compounds of formula I as claimed in claim 2 wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

R^4 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR^7R^8 , halogen or SH;

5 R^5 is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR^7R^8 , $NHOR^9$, $NHNR^7R^8$ or SH;

R^6 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR^7R^8 , halogen, SH or cyano;

10 R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

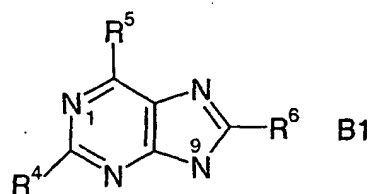
R^9 is hydrogen, alkyl or aryl;

with the proviso that R^4 is not NH_2 and R^5 is not $NH(CH_3)$.

9. Use of compounds of formula I as claimed in any one of claims 2 or 8

15 wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

R^4 is hydrogen or chlorine;

20 R^5 is hydroxy, alkylthio, aryl, heterocycl, halogen, NR^7R^8 or SH;

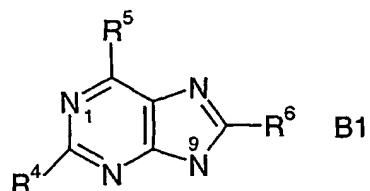
R^6 is hydrogen, halogen, heterocycl or NR^7R^8 ;

R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, alkenylalkyl or alkynylalkyl;

with the proviso that R⁵ is not NH(CH₃).

- 5 10. Use of compounds of formula I as claimed in any one of claims 2, 8 or 9
wherein

B signifies a purine base B1 which is connected through the 9-nitrogen of formula



wherein

10 R⁴ is hydrogen;

R⁵ is alkylthio, aryl, heterocyclyl, halogen or NR⁷R⁸;

R⁶ is hydrogen or halogen;

R⁷ and R⁸ are independently of each other hydrogen, alkyl, alkenylalkyl or alkynylalkyl;

15 with the proviso that R⁵ is not NH(CH₃).

11. Use of a compound of formula I as claimed in claim 1 which compound is

6-Dimethylamino-9-(β-D-ribofuranosyl)purine,

6-[1(S)-Methyl-2-phenylethylamino]-9-(β-D-ribofuranosyl)purine,

20 3'-Deoxyadenosine,

- 6-(Phenylethylamino)-9-(β -D-ribofuranosyl)purine,
6-(Cyclohexylamino)-9-(β -D-ribofuranosyl)purine,
2-Chloroadenosine,
9-(β -D-Ribofuranosyl)purine,
5 8-Bromoadenosine,
8-Bromo-2'-deoxyadenosine,
8-Bromoguanosine,
6-Thioinosine,
6-Methylthio-9-(β -D-ribofuranosyl)purine,
10 6-Chloro-9-(β -D-ribofuranosyl)purine,
2-Amino-6-chloro-9-(β -D-ribofuranosyl)purine,
6-(N-Methylpropylamino)-9-(β -D-ribofuranosyl)purine,
9-(β -D-Ribofuranosyl)-6-(4-thiomorpholinyl)purine,
6-(N-Methyl-2-propenylamino)-9-(β -D-ribofuranosyl)purine,
15 6-(N-Methyl-2-propynylamino)-9-(β -D-ribofuranosyl)purine,
6-(4-Morpholinyl)-9-(β -D-ribofuranosyl)purine,
6-Diethylamino-9-(β -D-ribofuranosyl)purine,
6-(1(R,S)-Phenylethylamino)-9-(β -D-ribofuranosyl)purine,
6-(1-Benzyl-1-methylethylamino)-9-(β -D-ribofuranosyl)purine,
20 6-(3-Phenylpropylamino)-9-(β -D-ribofuranosyl)purine,

- 9-(β -D-Ribofuranosyl)-6-[2-(2-thienyl)ethylamino]purine,
- 6-Dibenzylamino-9-(β -D-ribofuranosyl)purine,
- 6-Hexylamino-9-(β -D-ribofuranosyl)purine,
- 6-(3-Pyridylmethylamino)-9-(β -D-ribofuranosyl)purine,
- 5 6-[4-(4-Fluorophenyl)-1,2,5,6-tetrahydropyridyl]-9-(β -D-ribofuranosyl)purine,
- 6-[4-(2-Methoxyphenyl)piperazinyl]-9-(β -D-ribofuranosyl)purine,
- 6-[2-(3-Indolyl)ethylamino]-9-(β -D-ribofuranosyl)purine,
- 6-[2-(4-Chlorophenyl)ethylamino]-9-(β -D-ribofuranosyl)purine,
- 6-(N-Methylphenylamino)-9-(β -D-ribofuranosyl)purine,
- 10 9-(β -D-Ribofuranosyl)-6-(1,2,4,5-tetrahydro-3H-benzazepin-3-yl)purine,
- 9-(β -D-Ribofuranosyl)-6-(1,2,3,4-tetrahydro-2-isoquinolyl)purine,
- 6-(4-Methylpiperazinyl)-9-(β -D-ribofuranosyl)purine,
- 9-(β -D-Ribofuranosyl)-6-(1,3,4,5-tetrahydro-2H-benzazepin-2-yl)purine,
- 6-[2-(4-Cyanomethylphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine,
- 15 6-(2,3-Dihydro-1-indolyl)-9-(β -D-ribofuranosyl)purine,
- 9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl)purine,
- 9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl)purine,
- 6-(8-Aminosulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl)-9-(β -D-ribofuranosyl)purine,
- 20 6-[2-(3,4-Dimethoxyphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine,

- 6-[2-(4-Hydroxyphenyl)ethylamino]-9-(β -D-ribofuranosyl)purine,
- 6-(2-Isoindolinyl)-9-(β -D-ribofuranosyl)purine,
- 6-(7-Aminosulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl)-9-(β -D-ribofuranosyl)purine,
- 5 6-(N-Cyclohexylmethylamino)-9-(β -D-ribofuranosyl)purine,
- 6-(N-Hexylmethylamino)-9-(β -D-ribofuranosyl)purine,
- 6-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-ylamino)-9-(β -D-ribofuranosyl)purine,
- 10 6-[N-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-yl)methylamino]-9-(β -D-ribofuranosyl)purine,
- 6-[N-(5-Aminopentyl)methylamino]-9-(β -D-ribofuranosyl)purine,
- 6-[(5-Chloro-2-methoxyphenyl)methylamino]-9-(β -D-ribofuranosyl)purine,
- 6-[(2-Methylphenyl)methylamino]-9-(β -D-ribofuranosyl)purine,
- 6-(Hexamethyleneimino)-9-(β -D-ribofuranosyl)purine,
- 15 6-(1-Pyrrolidinyl)-9-(β -D-ribofuranosyl)purine,
- 6-(4-Hydroxypiperidin-1-yl)-9-(β -D-ribofuranosyl)purine,
- 6-(1-Piperidinyl)-9-(β -D-ribofuranosyl)purine,
- 6-(2-Propenyl)amino-9-(β -D-ribofuranosyl)purine,
- 6-(2-Propynyl)amino-9-(β -D-ribofuranosyl)purine,
- 20 6-(1-Methyl)ethylamino-9-(β -D-ribofuranosyl)purine,
- 6-bis-(2-Propenyl)amino-9-(β -D-ribofuranosyl)purine,

- 6-(2-Phenylethyl)methylamino-9-(β -D-ribofuranosyl)purine,
- 6-Ethylmethylamino- 9-(β -D-ribofuranosyl)purine,
- 6-bis-[(3-Methyl)butylamino]-9-(β -D-ribofuranosyl)purine,
- 6-(4-Aminophenyl)methylamino-9-(β -D-ribofuranosyl)purine,
- 5 6-(2-Pyridylmethyl)amino-9-(β -D-ribofuranosyl)purine,
- 6-(2-Hydroxyethyl)methylamino-9-(β -D-ribofuranosyl)purine,
- 6-Dipropylamino-9-(β -D-ribofuranosyl)purine,
- 6-[2-Phenyl-(N-propionyl)ethylamino]-9-(β -D-ribofuranosyl)purine,
- 6-(N-Benzoyl-2-phenylethylamino)-9-(β -D-ribofuranosyl)purine,
- 10 2-Amino-6-methylamino-9-(β -L-ribofuranosyl)purine,
- 2-Amino-6-methylamino-9-(β -D-ribofuranosyl)purine,
- 2-Amino-6-(4-morpholinyl)-9-(β -D-ribofuranosyl)purine,
- 2-Amino-6-(1-pyrrolidinyl)-9-(β -D-ribofuranosyl)purine,
- 2,6-Diamino-9-(β -L-ribofuranosyl)purine,
- 15 2,6-Diamino-9-(β -D-ribofuranosyl)purine,
- 2-Chloro-6-(1-pyrrolidinyl)-9-(β -D-ribofuranosyl)purine,
- 2-Chloro-6-(1-hexamethyleneimino)-9-(β -D-ribofuranosyl)purine,
- 2-Chloro-6-(4-hydroxy-1-piperidinyl)-9-(β -D-ribofuranosyl)purine,
- 6-[(N-Cyclohexyl)methylamino]-2-methylthio-9-(β -D-ribofuranosyl)purine,
- 20 6-(1-Pyrrolyl)-9-(β -D-ribofuranosyl)purine,

- 6-(1-Pyrrolyl)-9-(β -D-arabinofuranosyl)purine,
- 6-(1-Pyrrolyl)-9-(β -D-ribofuranosyl)purin-8-(7H)-one,
- 9-(3-Deoxy- β -D-ribofuranosyl)-6-(1-pyrrolyl) purine,
- 6-(1-Pyrrolyl)-9-(β -L-ribofuranosyl)purine,
- 5 6-(1-Indolyl)-9-(β -D-ribofuranosyl)purine,
- 6-(1-Imidazolyl)-9-(β -D-ribofuranosyl)purine,
- 9-(β -D-Ribofuranosyl)-6-(1,2,4-triazol-1-yl)purine,
- 6-(1-Pyrazolyl)- 9-(β -D-ribofuranosyl)purine,
- 9-(β -D-Ribofuranosyl) 6-(1,2,4-triazol-4-yl)purine,
- 10 6-Methylamino-9-(β -D-ribofuranosyl)purin-2(1H)-one,
- 2-Methoxy-6-methylamino-9-(β -D-ribofuranosyl)purine,
- 2-Methoxyadenosine,
- 2,6-Dichloro-9-(β -D-ribofuranosyl)purine,
- 6-Methoxy-9-(β -D-ribofuranosyl)purine,
- 15 2-Amino-6-benzylthio-9-(β -D-ribofuranosyl)purine,
- 6-Benzylthio-2-hydroxy-9-(β -D-ribofuranosyl)purine,
- 9-(β -D-Ribofuranosyl)purine-2,6,8(1H,3H,7H)-trione,
- 8-(Methylamino)adenosine,
- 8-(2-Phenylethylamino)adenosine,
- 20 8-Benzylaminoadenosine,

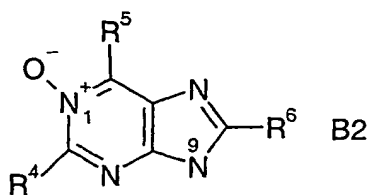
- 8-(1-Piperidinyl)adenosine,
8-(Dimethylamino)adenosine,
8-(3-Phenylpropylamino)adenosine,
8-(4-Morpholinyl)adenosine,
5 8-(N-Methyl-2-phenylethylamino)adenosine,
8-(3-Pyridylmethylamino)adenosine,
8-(Ethylamino)adenosine,
8-(1,2,3,4-Tetrahydro-2-isoquinolyl)adenosine,
8-[2-(4-Morpholinyl)ethylamino]adenosine,
10 8-(Hexylamino)adenosine,
8-(2-Cyclohexylethylamino)adenosine,
8-(2(R,S)-Phenylpropylamino)adenosine,
8-[2-(4-Methylphenyl) ethylamino]adenosine,
8-[2-(1-methyl-2-pyrrolyl) ethylamino]adenosine,
15 8-[2-(4-Aminosulphonylphenyl) ethylamino]adenosine,
8-(4-Phenyl-1-piperazinyl)adenosine,
8-(2-(4-Imidazolyl)adenosine,
8-(1-Naphthylmethylamino)adenosine,
8-[2-(4-Hydroxyphenyl)ethylamino]adenosine,
20 8-(4-Phenylbutylamino)adenosine,
8-[2-(4-Chlorophenyl)ethylamino]adenosine,
8-[2-(2,4-Dichlorophenyl)ethylamino]adenosine,

- 8-(2-Propenylamino)adenosine,
- 8-(2-Hydroxyethylamino)adenosine,
- 8-(1(R)-Methyl-2-phenylethylamino)adenosine,
- 8-(4-Fluorobenzylamino)adenosine,
- 5 8-[(4-Hydroxycarbonyl)benzylamino]adenosine,
- 8-(2-Propynylamino)adenosine,
- 8-(1-Methylethylamino)adenosine,
- 8-[(4-Trifluoromethyl)benzylamino]adenosine,
- 8-[(2,5-Dimethoxy)benzylamino]adenosine,
- 10 8-[2-(2-Thienyl)ethylamino]adenosine,
- 8-[2-(4-Aminophenyl)ethylamino]adenosine,
- 8-(2-Phenoxyethylamino)adenosine,
- 8-[(2-Thienyl)methylamino]adenosine,
- 8-[(4-tert-Butyl)benzylamino]adenosine,
- 15 8-(1(R)-Phenylethylamino)adenosine,
- 8-(1(S)-Phenylethylamino)adenosine,
- 8-(6-Phenylhexylamino)adenosine,
- 8-[2-Hydroxy-1(S)-phenyl)ethylamino]adenosine,
- 2'-Deoxy-8-(2-phenylethylamino)adenosine,
- 20 2'-Deoxy-8-(3-phenylpropylamino)adenosine,
- 8-Benzylamino-2'-deoxyadenosine,
- 2'-Deoxy-8-(4-phenylbutylamino)adenosine,

- 2'-Deoxy-8-(6-phenylhexylamino)adenosine,
8-(4-Morpholinyl)inosine,
8-(Methylthio)adenosine,
8-(Benzylthio)adenosine,
5 8-(Benzyoxy)adenosine,
8-Ethoxyadenosine,
8-[(1-Hydroxy-1-methyl)ethyl]adenosine,
9-(β -D-ribofuranosyl)-6-(3-thienyl)purine,
6-Phenyl-9-(β -D-ribofuranosyl) purine,
10 6-(4-Fluorophenyl)-9-(β -D-ribofuranosyl) purine,
6-(4-Chlorophenyl)-9-(β -D-ribofuranosyl) purine,
6-(4-Methylphenyl)-9-(β -D-ribofuranosyl) purine,
6-(4-Methoxyphenyl)-9-(β -D-ribofuranosyl) purine,
9-(β -D-Ribofuranosyl)-6-(1-thianthrenyl)purine,
15 6-(4-Biphenylyl)-9-(β -D-ribofuranosyl) purine,
6-(4-Methylthiophenyl)-9-(β -D-ribofuranosyl) purine,
6-(2-Methylphenyl)-9-(β -D-ribofuranosyl) purine,
6-(9-Phenanthrenyl)-9-(β -D-ribofuranosyl)purine,
9-(β -D-Ribofuranosyl)-6-(3-trifluoromethylphenyl)purine,
20 6-(2-Phenoxyphenyl)-9-(β -D-ribofuranosyl) purine,

- 6-(4-tert-Butylphenyl)-9-(β -D-ribofuranosyl) purine,
- 9-(β -D-Ribofuranosyl)-6-(2-trifluoromethoxyphenyl)purine,
- 6-(4-Phenoxyphenyl)-9-(β -D-ribofuranosyl)purine,
- 6-(3-Methoxyphenyl)-9-(β -D-ribofuranosyl) purine,
- 5 6-(2-Naphthyl)-9-(β -D-ribofuranosyl)purine,
- 6-(3-Biphenylyl)-9-(β -D-ribofuranosyl)purine,
- 6-[4-(2-Methylpropyl)phenyl]-9-(β -D-ribofuranosyl)purine,
- 6-(3-Fluorophenyl)-9-(β -D-ribofuranosyl)purine,
- 9-(β -D-Ribofuranosyl)-6-(4-trifluoromethylphenyl)purine,
- 10 6-(3-Ethoxyphenyl)-9-(β -D-ribofuranosyl)purine,
- 6-[3-(1-Methyl)ethylphenyl]-9-(β -D-ribofuranosyl)purine,
- 9-(β -D-ribofuranosyl)-6-(4-trifluoromethoxyphenyl)purine,
- 6-(4-Ethylphenyl)-9-(β -D-ribofuranosyl)purine,
- 2-Amino-6-phenyl-9-(β -D-ribofuranosyl)purine,
- 15 6-Ethylamino-9-(β -D-ribofuranosyl)purine, or
- 6-Propylamino-9-(β -D-ribofuranosyl)purine.

12. Use of compounds of formula I as claimed in any one of claims 1 to 4
wherein
- 20 B signifies an oxidised purine base B2 which is connected through the 9-nitrogen
of formula



wherein

R^4 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen or SH;

5 R^5 is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclylamino, halogen, NR^7R^8 , $NHOR^9$, $NHNR^7R^8$ or SH;

R^6 is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl, NR^7R^8 , halogen, SH or cyano;

10 R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R^9 is hydrogen, alkyl or aryl.

13. Use of compounds of formula I as claimed in any one of claims 1 to 4 or 12

15 wherein

R^4 is hydrogen;

R^5 is hydrogen, alkyl, heterocyclyl or NR^7R^8 ;

R^6 is hydrogen;

20 R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl.

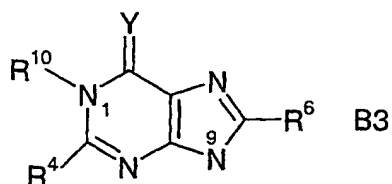
14. Use of a compound of formula I as claimed in any one of claims 1 to 4 or 12 to 13 which compound is

Adenosine-1-oxide, or

6-(2-Phenylethylamino)- 9-(β -D-ribofuranosyl)purine-1-oxide.

15. Use of compounds of formula I as claimed in any one of claims 1 to 4
5 wherein

B signifies a purine base B3 which is connected through the 9-nitrogen of formula



wherein

R⁴ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl,
NR⁷R⁸, halogen or SH;

R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocyclyl,
NR⁷R⁸, halogen, SH or cyano;

R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl,
alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

15 R⁹ is hydrogen, alkyl or aryl;

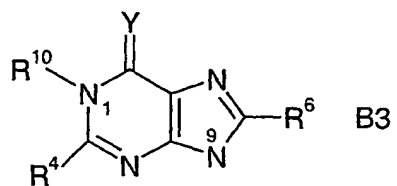
R¹⁰ is hydrogen, alkyl or aryl;

Y is O, S or NR¹¹;

R¹¹ is hydrogen, hydroxy, alkyl, OR⁹, heterocyclyl or NR⁷R⁸.

- 20 16. Use of compounds of formula I as claimed in any one of claims 1 to 4 or 15
wherein

B signifies a purine base B3 which is connected through the 9-nitrogen of formula



R⁴ is hydrogen, NR⁷R⁸ or hydroxy;

R⁶ is hydrogen, halogen or NR⁷R⁸;

5 R⁷ and R⁸ are independently of each other hydrogen or alkyl;

R¹⁰ is hydrogen or alkyl;

Y is O, S, NH or N-alkyl.

17. Use of a compound of formula I as claimed in claim 1 which compound is

10 3'-Deoxyguanosine,

6-Thioguanosine,

Inosine,

L-Inosine,

8-Bromoinosine,

15 1-Benzyl-6-imino-9-(β-D-ribofuranosyl)purine,

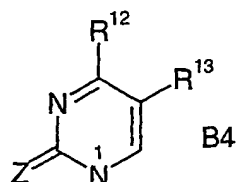
1-Methyl-6-(2-phenylethylimino)-9-(β-D-ribofuranosyl)purine,

2-(Acetylamino)inosine, or

8-(Benzylamino)inosine.

18. Use of compounds of formula I as claimed in any one of claims 1, 3 or 4
wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



5

wherein

Z is O or S;

R¹² is hydrogen, hydroxy, alkyl, alkoxy, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH;

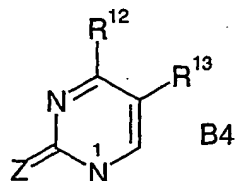
10 R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen;

R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R⁹ is hydrogen, alkyl or aryl.

- 15 19. Use of compounds of formula I as claimed in any one of claims 1, 3, 4 or 18
wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



20

wherein

Z is O;

R¹² is hydroxy, alkyl, heterocyclyl, NR⁷R⁸, NHOR⁹, heterocyclylamino, NHNR⁷R⁸ or SH;

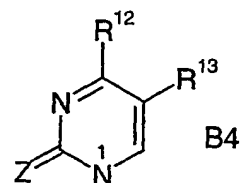
R¹³ is hydrogen, alkyl or halogen;

5 R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R⁹ is hydrogen, alkyl or aryl.

20. Use of compounds of formula I as claimed in any one of claims 1, 3, 4, 18 or
10 19 wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

15 Z is O;

R¹² is hydroxy, alkyl or NR⁷R⁸;

R¹³ is hydrogen;

R⁷ and R⁸ are independently of each other hydrogen or alkyl.

20 21. Use of compounds of formula I as claimed in claim 1 wherein

R¹ is hydrogen, halogen, hydroxy, alkyl, alkoxy, cyano or azido;

R² is hydrogen or hydroxy; or

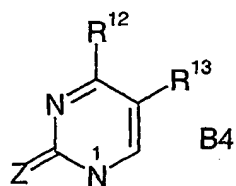
R² and R³ represent fluorine;

X is O or CH₂;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

5

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

10 Z is O;

R¹² is NR⁷R⁸;

R¹³ is hydrogen, alkyl or halogen;

R⁷ and R⁸ are independently of each other hydrogen or alkyl.

15 22. Use of compounds of formula I as claimed in claim 1 or 21 wherein

R¹ is hydrogen, fluorine, hydroxy, C₁₋₄-alkyl, C₁₋₄-alkoxy, cyano or azido;

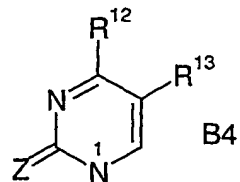
R² is hydrogen or hydroxy; or

R² and R³ represent fluorine;

X is O or CH₂;

20 a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

5 Z is O;

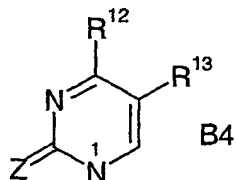
R¹² is NR⁷R⁸;

R¹³ is hydrogen, C₁₋₄-alkyl or fluorine;

R⁷ and R⁸ are independently of each other hydrogen or C₁₋₄-alkyl.

10 23. Use of compounds of formula I as claimed in claim 2 wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

15 Z is O or S;

R¹² is hydrogen, alkyl, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocyclyl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH;

R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen;

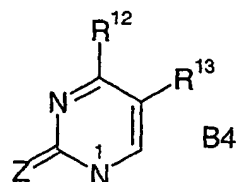
R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R^9 is hydrogen, alkyl or aryl;

with the proviso that R^{12} is not $N(CH_3)_2$, $N(H)NH(CH_3)$ or $N(H)NH_2$ and R^{13} is
5 not hydroxyalkyl, chlorine or bromine.

24. Use of compounds of formula I as claimed in any one of claims 2 or 23
wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of
10 formula



wherein

Z is O;

R^{12} is alkyl, heterocyclyl, NR^7R^8 , $NHOR^9$, heterocyclylamino, $NHNR^7R^8$ or SH;

15 R^{13} is hydrogen, alkyl or halogen;

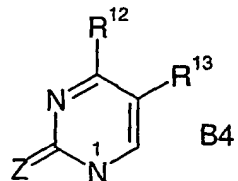
R^7 and R^8 are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl;

R^9 is hydrogen, alkyl or aryl;

with the proviso that R^{12} is not $N(CH_3)_2$, $N(H)NH(CH_3)$ or $N(H)NH_2$ and R^{13} is
20 not hydroxyalkyl, chlorine or bromine.

25. Use of compounds of formula I as claimed in any one of claims 2, 23 or 24
wherein

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of
formula



5

wherein

Z is O;

R¹² is alkyl or NR⁷R⁸;

R¹³ is hydrogen;

10 R⁷ and R⁸ are independently of each other hydrogen or alkyl;

with the proviso that R¹² is not N(CH₃)₂, N(H)NH(CH₃) or N(H)NH₂.

26. Use of a compound of formula I as claimed in claim 1 which compound is

4-Thiouridine,

15 5-Fluorocytidine,

1-(β-D-arabinofuranosyl)-5-fluorocytosine,

5-Methylcytidine,

2',3'-Dideoxycytidine,

N4-Acetylcytidine,

20 3'-Deoxycytidine,

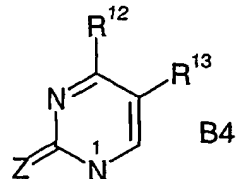
- 4-Methoxy-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
4-Methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
5-Fluoro-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
5-Methyl-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
5
3'-Azido-2',3'-dideoxy-5-methylcytidine,
1-(3-Deoxy- β -L-threo-pentofuranosyl)-5-fluorocytosine,
4-Methylamino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
5-Fluoro-4-methylamino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
4-(1-Pyrrolyl)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,
10
1-(2-Deoxy-2,2-difluoro- β -D-erythropentofuranosyl)cytosine,
4-Amino-1(R)-(2(S),3(R)-dihydroxy-4(R)-hydroxymethyl-cyclopentyl)-1H-pyrimidin-2-one,
1-(β -D-Xylofuranosyl)cytosine,
1-(3-Deoxy-3-fluoro- β -D-xylofuranosyl)cytosine, or
15
3'-Deoxy-3'-hydroxymethylcytidine.

27. Use of compounds of formula I as claimed in claim 2 wherein

- R¹ is hydrogen, halogen, hydroxy, alkyl, alkoxy, cyano or azido;
R² is hydrogen or hydroxy; or
20 R² and R³ represent fluorine;
X is O or CH₂;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



5

wherein

Z is O;

R¹² is NR⁷R⁸;

R¹³ is hydrogen, alkyl or halogen;

10 R⁷ and R⁸ are independently of each other hydrogen or alkyl;

with the proviso that R¹² is not N(CH₃)₂ and R¹³ is not chlorine or bromine.

28. Use of compounds of formula I as claimed in claim 2 or 27 wherein

R¹ is hydrogen, fluorine, hydroxy, C₁₋₄-alkyl, C₁₋₄-alkoxy, cyano or azido;

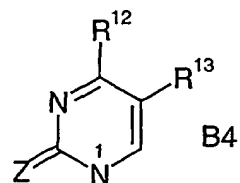
15 R² is hydrogen or hydroxy; or

R² and R³ represent fluorine;

X is O or CH₂;

a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

20 B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula



wherein

Z is O;

R¹² is NR⁷R⁸;

5 R¹³ is hydrogen, C₁₋₄-alkyl or fluorine;

R⁷ and R⁸ are independently of each other hydrogen or C₁₋₄-alkyl;

with the proviso that R¹² is not N(CH₃)₂.

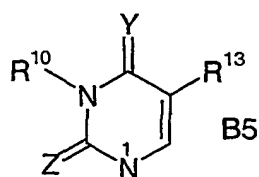
29. Use of a compound of formula I as claimed in any one of claims 1, 21, 22, 27
10 or 28 which compound is

L-Cytidine, or

4-Amino-1-(2,2-difluoro-3-hydroxy-4-hydroxymethyl-cyclopentyl)-1H-pyrimidin-2-one.

15 30. Use of compounds of formula I as claimed in any one of claims 1, 3 or 4
 wherein

B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of
formula

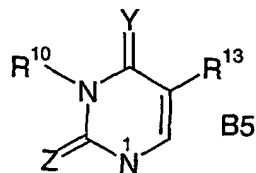


wherein

- Y is O, S or NR¹¹;
- Z is O or S;
- R¹⁰ is hydrogen, alkyl or aryl;
- 5 R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen.

31. Use of compounds of formula I as claimed in any one of claims 1 to 4 or 30
wherein

- B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of
10 formula

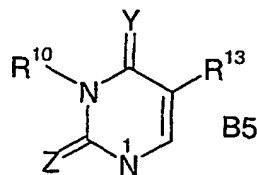


wherein

- Y is O or NR¹¹;
- Z is O;
- 15 R¹⁰ is hydrogen;
- R¹³ is hydrogen, alkyl or halogen.

32. Use of compounds of formula I as claimed in claim 2 wherein

- B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of
20 formula



wherein

Y is O, S or NR¹¹;

Z is O or S;

5 R¹⁰ is hydrogen, alkyl or aryl;

R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen;

with the proviso that R¹⁰ is not methyl or hydroxyethyl.

33. Use of a compound of formula I as claimed in claim 1 which compound is

10 2'-Deoxy-5-fluorouridine,

1-(β-D-Arabinofuranosyl) -5-fluorouracil,

5-Fluorouridine,

5-Bromouridine,

3-Methyluridine,

15 5-Methyluridine,

1-(β-D-Arabinofuranosyl)uracil,

1-(β-D-Arabinofuranosyl)-5-methyluracil,

1-(β-D-Arabinofuranosyl)-5-iodouracil,

3'-Deoxy-5-methyluridine,

20 5-Ethyluridine,

- 5-[(1-Methyl)ethyl]uridine,
5-Methoxymethyluridine,
5-Ethoxymethyluridine,
5-Chlorouridine,
5 5-Methyl-1-(β -L-ribofuranosyl)uracil,
1-(β -D-Arabinofuranosyl)-5-ethyluracil,
1- (β -D-Arabinofuranosyl)-5-bromo uracil,
5-Methyl-4-thiouridine,
5-Fluoro-4-thiouridine,
10 1-(2-Deoxy - α -D-erthyro-pentofuranosyl)-5-fluorouracil,
2'-Deoxy-5-fluoro-3-methyluridine,
1-(α -D-Erthyro-2-deoxypentofuranosyl)-5-fluoro-3-methyluracil,
2'-Chloro-2'-deoxyuridine,
2'-Bromo-2'-deoxyuridine,
15 1-(2-Deoxy- β -D-lyxofuranosyl)-5-methyluracil,
3'-Deoxy-3'-fluoro-5-methyluridine,
2',3'-Dideoxy-5-ethyl-3'-methoxyuridine,
5'-Benzylxy-2',3'-dideoxy-5-methyluridine,
2',3'-Dideoxy-5-ethyl-3'-iodouridine,
20 3'-Azido-2',3'-dideoxy-5-ethyluridine,
4-Oximino-1-(β -L-ribofuranosyl)pyrimidin-2(1H)-one,

4-Oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,

4-Oximino-1-(β -D-arabinofuranosyl)pyrimidin-2(1H)-one,

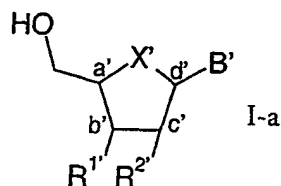
5-Fluoro-4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,

1-(2-Deoxy-2,2-difluoro- α -D-erythropentofuranosyl)uracil,

5 1-(3-Deoxy-3-fluoro- β -D-xylofuranosyl)uracil, or

2'-Deoxy-2'-methoxyuridine.

34. Compounds of formula I-a



10 wherein

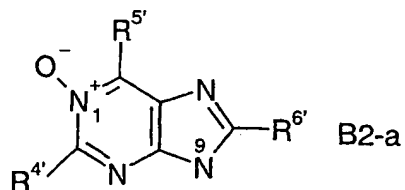
$R^{1\prime}$ is hydroxy;

$R^{2\prime}$ is hydroxy;

X' is O;

15 a', b', c', d' denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

B' signifies an oxidised purine base $B2-a$ which is connected through the 9-nitrogen of formula



wherein

R^{4'} is hydrogen;

R^{5'} is NHR^{8'};

5 R^{6'} is hydrogen;

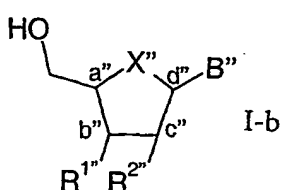
R^{8'} is alkyl;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

35. A compound as claimed in claim 34 which compounds are selected from:

10 6-(2-phenylethylamino)- 9-(β -D-ribofuranosyl)purine-1-oxide.

36. Compounds of formula I-b



wherein

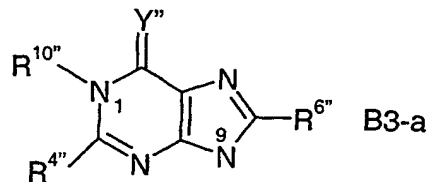
15 R^{1''} is hydroxy;

R^{2''} is hydroxy;

X'' is O;

a'' , b'' , c'' , d'' denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

B'' signifies a purine base B3-a which is connected through the 9-nitrogen of formula



5

wherein

$R^{4''}$ is hydrogen;

$R^{6''}$ is hydrogen;

$R^{10''}$ is alkyl;

10 Y'' is $NR^{11''}$;

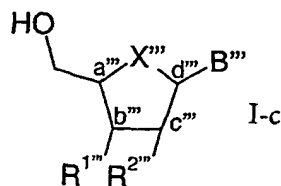
$R^{11''}$ is alkyl;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

37. A compound as claimed in claim 36 which compound is selected from:

15 1-Methyl-6-(2-phenylethylimino)-9-(β -D-ribofuranosyl)purine.

38. Compounds of formula I-c



wherein

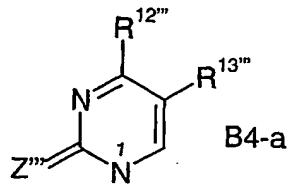
$R^{1''''}$ is hydroxy;

$R^{2''''}$ is hydroxy;

X''' is O;

5 a'''' , b'''' , c'''' , d'''' denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

group B'''' signifies a pyrimidine base B4-a which is connected through the 1-nitrogen of formula



10 wherein

$R^{12''''}$ is alkylthio or heterocyclyl;

$R^{13''''}$ is hydrogen, alkyl or halogen;

Z''' is O;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

15

39. Compound as claimed in claim 38 which compound is

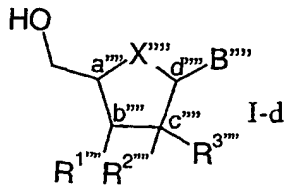
5-Fluoro-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,

5-Methyl-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one, or

4-(1-Pyrrolyl)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one.

20

40. Compounds of formula I-d



wherein

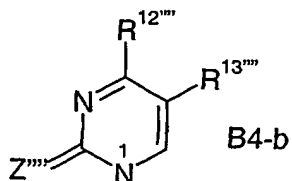
$R^{1'''}$ is hydrogen, halogen, hydroxy, alkyl, alkoxy, cyano or azido;

5 $R^{2'''}$ and $R^{3'''}$ represent fluorine;

X''' is O or CH_2 ;

a''', b''', c''', d''' denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and

10 group B''' signifies a pyrimidine base B4-b which is connected through the 1-nitrogen of formula



wherein

Z''' is O;

$R^{12'''}$ is $NR^{7'''}R^{8'''}$;

15 $R^{13'''}$ is hydrogen, alkyl or halogen;

$R^{7'''}$ and $R^{8'''}$ are independently of each other hydrogen or alkyl;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

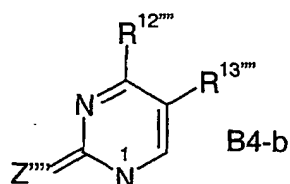
41. Compounds of formula I-d as claimed in claim 40

wherein

$R^{1''''}$ is hydrogen, fluorine, hydroxy, C_{1-4} -alkyl, C_{1-4} -alkoxy, cyano or azido;

X'''' is CH_2 ; and

- 5 group B'''' signifies a pyrimidine base B4-b which is connected through the 1-nitrogen of formula



wherein

$R^{12''''}$ is hydrogen, C_{1-4} -alkyl or fluorine;

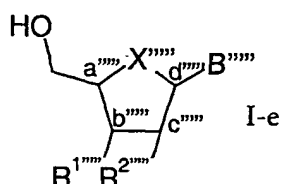
- 10 $R^{7''''}$ and $R^{8''''}$ are independently of each other hydrogen or C_{1-4} -alkyl.

42. Compound as claimed in claims 40 or 41 which compound is

4-Amino-1-(2,2-difluoro-3-hydroxy-4-hydroxymethyl-cyclopentyl)-1H-pyrimidin-2-one.

15

43. Compounds of formula I-e



wherein

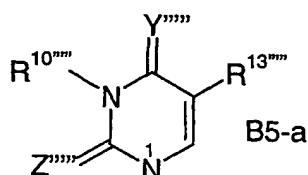
R^{1”””} is alkoxy;

R^{2”””} is hydrogen;

X””” is O;

5 a”””, b”””, c”””, d””” denoting asymmetric carbon atoms and forming a D-ribofuranosyl ring; and

group B””” signifies a pyrimidine base B5-a which is connected through the 1-nitrogen of formula



wherein

10 R^{10”””} is hydrogen;

R^{13”””} is alkyl;

Y””” is O;

Z””” is O;

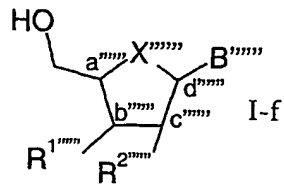
hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

15

44. Compound as claimed in claim 43 which compound is

2’,3’-Dideoxy-5-ethyl-3’-methoxyuridine.

45. Compounds of formula I-f



wherein

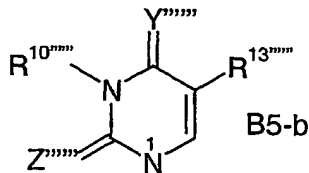
$R^{1''''''}$ is hydroxy;

$R^{2''''''}$ is hydroxy;

5 X'''''' is O;

a'''''' , b'''''' , c'''''' , d'''''' denoting asymmetric carbon atoms and forming a *D*-ribofuranosyl ring; and

group B'''''' signifies a pyrimidine base B5-b which is connected through the 1-nitrogen of formula



10

wherein

$R^{10''''''}$ is hydrogen;

$R^{13''''''}$ is halogen;

Y'''''' is $NR^{11''''''}$;

15 $R^{11''''''}$ is hydroxy;

Z'''''' is O;

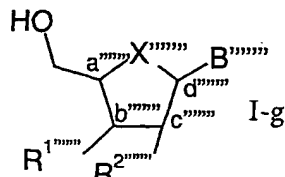
hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

46. Compound as claimed in claim 45 which compound is

5-Fluoro-4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one.

47. Compounds of formula I-g

5



wherein

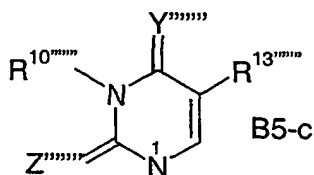
$R^{1''''''}$ is hydroxy;

$R^{2''''''}$ is hydroxy;

X'''''' is O;

10 $a'''''', b''''', c''''', d'''''$ denoting asymmetric carbon atoms and forming a L-ribofuranosyl ring; and

group B'''''' signifies a pyrimidine base B5-c which is connected through the 1-nitrogen of formula



15 wherein

$R^{10''''''}$ is hydrogen;

$R^{13''''''}$ is hydrogen;

Y'''''' is $NR^{11''''''}$;

R¹¹ is hydroxy;

Z is O;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

5 48. Compound as claimed in claim 47 which compound is

4-Oximino-1-(β-L-ribofuranosyl)pyrimidin-2(1H)-one.

49. Compound of formula I as claimed in claim 1 which compound is

6-(N-Methylpropylamino)-9-(β-D-ribofuranosyl)purine,

10 9-(β-D-Ribofuranosyl)-6-(4-thiomorpholinyl)purine,

6-(N-(2-Propenyl)methylamino)-9-(β-D-ribofuranosyl)purine,

6-(N-Methyl-2-propynylamino)-9-(β-D-ribofuranosyl)purine,

6-[4-(4-Fluorophenyl)-1,2,5,6-tetrahydropyridyl]-9-(β-D-ribofuranosyl)purine,

6-[4-(2-Methoxyphenyl)piperazinyl]-9-(β-D-ribofuranosyl)purine,

15 6-(N-Methylphenylamino)-9-(β-D-ribofuranosyl)purine,

9-(β-D-Ribofuranosyl)-6-(1,2,4,5-6-(1-itetrahydro-3H-benzazepin-3-yl)purine,

9-(β-D-ribofuranosyl)-6-(1,2,3,4-tetrahydro-2-isoquinolyl)purine,

9-(β-D-Ribofuranosyl)-6-(1,3,4,5-tetrahydro-2H-benzazepin-2-yl)purine,

6-[2-(4-Cyanomethylphenyl)ethylamino]-9-(β-D-ribofuranosyl)purine,

20 6-(2,3-Dihydro-1-indolyl)- 9-(β-D-ribofuranosyl)purine,

- 9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzothiazepin-4-yl)purine,
- 9-(β -D-Ribofuranosyl)-6-(2,3,4,5-tetrahydro-1,4-benzoxazepin-4-yl)purine,
- 6-(8-Aminosulphonyl-2,3,4,5-tetrahydro-1H-2-benzazepin-2-yl)-9-(β -D-ribofuranosyl)purine,
- 5 6-(2-Isoindolinyl)-9-(β -D-ribofuranosyl)purine,
- 6-(7-Aminosulphonyl-2,3,4,5-tetrahydro-1H-benzazepin-3-yl)-9-(β -D-ribofuranosyl)purine,
- 6-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-ylamino)-9-(β -D-ribofuranosyl)purine,
- 10 6-[N-(10,11-Dihydro-5H-dibenzo[a,d]cyclohepten-5-yl)methylamino]-9-(β -D-ribofuranosyl)purine,
- 6-[N-(5-Aminopentyl)methylamino]-9-(β -D-ribofuranosyl)purine,
- 6-Ethylmethylamino- 9-(β -D-ribofuranosyl)purine,
- 6-bis-[(3-Methyl)butylamino]-9-(β -D-ribofuranosyl)purine,
- 15 6-[2-Phenyl-(N-propionyl)ethylamino]-9-(β -D-ribofuranosyl)purine,
- 6-(N-Benzoyl-2-phenylethylamino)-9-(β -D-ribofuranosyl)purine,
- 1-Methyl-6-(2-phenylethylimino)-9-(β -D-ribofuranosyl)purine,
- 2-Amino-6-methylamino-9-(β -L-ribofuranosyl)purine,
- 6-[(N-Cyclohexyl)methylamino]-2-methylthio-9-(β -D-ribofuranosyl)purine,
- 20 6-(1-Pyrrolyl)-9-(β -D-ribofuranosyl)purin-8-(7H)-one,
- 9-(3-Deoxy- β -D-ribofuranosyl)-6-(1-pyrrolyl) purine,

- 6-(1-Pyrrolyl)-9-(β -L-ribofuranosyl)purine,
6-(1-Indolyl)-9-(β -D-ribofuranosyl)purine,
6-(1-Imidazolyl)-9-(β -D-ribofuranosyl)purine,
9-(β -D-Ribofuranosyl)-6-(1,2,4-triazol-1-yl)purine,
5 6-(1-Pyrazolyl)- 9-(β -D-ribofuranosyl)purine,
6-(2-Phenylethylamino)- 9-(β -D-ribofuranosyl)purine-1-oxide,
8-(2-Phenylethylamino)adenosine,
8-(3-Phenylpropylamino)adenosine,
8-(4-Morpholinyl)adenosine,
10 8-(N-Methyl-2-phenylethylamino)adenosine,
8-(3-Pyridylmethylamino)adenosine,
8-(1,2,3,4-Tetrahydro-2-isoquinolyl)adenosine,
8-[2-(4-Morpholinyl)ethylamino]adenosine,
8-(2-Cyclohexylethylamino)adenosine,
15 8-(2(R,S)-Phenylpropylamino)adenosine,
8-[2-(4-Methylphenyl) ethylamino]adenosine,
8-[2-(1-methyl-2-pyrrolyl) ethylamino]adenosine,
8-[2-(4-Aminosulphonylphenyl) ethylamino]adenosine,
8-(4-Phenyl-1-piperazinyl)adenosine,
20 8-(1-Naphthylmethylamino)adenosine,
8-[2-(4-Hydroxyphenyl)ethylamino]adenosine,

- 8-(4-Phenylbutylamino)adenosine,
- 8-[2-(4-Chlorophenyl)ethylamino]adenosine,
- 8-[2-(2,4-Dichlorophenyl)ethylamino]adenosine,
- 8-(2-Propenylamino)adenosine,
- 5 8-(1(R)-Methyl-2-phenylethylamino)adenosine,
- 8-(4-Fluorobenzylamino)adenosine,
- 8-[(4-Hydroxycarbonyl)benzylamino]adenosine,
- 8-(2-propynylamino)adenosine,
- 8-[(4-trifluoromethyl)benzylamino]adenosine,
- 10 8-[(2,5-Dimethoxy)benzylamino]adenosine,
- 8-[2-(2-Thienyl)ethylamino]adenosine,
- 8-[2-(4-Aminophenyl)ethylamino]adenosine,
- 8-(2-Phenoxyethylamino)adenosine,
- 8-[(2-Thienyl)methylamino]adenosine,
- 15 8-[(4-tert-Butyl)benzylamino]adenosine,
- 8-(1(R)-Phenylethylamino)adenosine,
- 8-(1(S)-Phenylethylamino)adenosine,
- 8-(6-Phenylhexylamino)adenosine,
- 8-[2-Hydroxy-1(S)-phenyl]ethylamino]adenosine,
- 20 2'-Deoxy-8-(2-phenylethylamino)adenosine,
- 2'-Deoxy-8-(3-phenylpropylamino)adenosine,
- 8-Benzylamino-2'-deoxyadenosine,

- 2'-Deoxy-8-(4-phenylbutylamino)adenosine,
2'-Deoxy-8-(6-phenylhexylamino)adenosine,
8-Ethoxyadenosine,
9-(β -D-Ribofuranosyl)-6-(3-thienyl)purine,
5 9-(β -D-Ribofuranosyl)-6-(1-thianthrenyl)purine,
6-(4-Biphenylyl)-9-(β -D-ribofuranosyl) purine,
6-(4-Methylthiophenyl)-9-(β -D-ribofuranosyl) purine,
6-(9-Phenanthrenyl)-9-(β -D-ribofuranosyl)purine,
9-(β -D-Ribofuranosyl)-6-(3-trifluoromethylphenyl)purine,
10 6-(2-Phenoxyphenyl)-9-(β -D-ribofuranosyl) purine,
6-(4-tert-Butylphenyl)-9-(β -D-ribofuranosyl) purine,
9-(β -D-Ribofuranosyl)-6-(2-trifluoromethoxyphenyl)purine,
6-(4-Phenoxyphenyl)-9-(β -D-ribofuranosyl)purine,
6-(2-Naphthyl)-9-(β -D-ribofuranosyl)purine,
15 6-(3-Biphenylyl)-9-(β -D-ribofuranosyl)purine,
6-[4-(2-Methylpropyl)phenyl]-9-(β -D-ribofuranosyl)purine,
6-(3-Fluorophenyl)-9-(β -D-ribofuranosyl)purine,
9-(β -D-Ribofuranosyl)-6-(4-trifluoromethylphenyl)purine,
6-(3-Ethoxyphenyl)-9-(β -D-ribofuranosyl)purine,
20 6-[3-(1-Methyl)ethylphenyl]-9-(β -D-ribofuranosyl)purine,

9-(β -D-Ribofuranosyl)-6-(4-trifluoromethoxyphenyl)purine,

6-(4-Ethylphenyl)-9-(β -D-ribofuranosyl)purine,

5-Fluoro-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,

5-Methyl-4-methylthio-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,

5 2',3'-Dideoxy-5-ethyl-3'-methoxyuridine,

4-(1-Pyrrolyl)-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one,

4-Oximino-1-(β -L-ribofuranosyl)pyrimidin-2(1H)-one, or

5-Fluoro-4-oximino-1-(β -D-ribofuranosyl)pyrimidin-2(1H)-one;

hydrolyzable esters or ethers thereof and pharmaceutically acceptable salts thereof.

10

50. A compound, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in any one of claims 34 to 49 for use in the treatment of a human or animal body.

15

51. A compound, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in claim 50 for its use in the treatment of Hepatitis C Virus (HCV) infections.

20

52. Use of compounds, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in any one of claims 34 to 49 for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment.

53. A pharmaceutical composition comprising a pharmaceutically effective amount of a compound, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in any one of claims 34 to 49 and, if desired, a pharmaceutical inert carrier, especially for use in the treatment of an Hepatitis C Virus (HCV) infection.

5

54. A process for preparing a medicament which process comprises bringing a compound, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in any one of claims 34 to 49 into a galenical administration form together with a pharmaceutical inert carrier.

10

55. A method of treating an Hepatitis C Virus (HCV) infection in a subject, which method comprises administering to said subject a therapeutically effective amount of a compound of formula I as defined in any one of claims 1 to 33, or a compound, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in any one of claims 34 to 49.

15

56. Use of compounds of formula I as defined in any one of claims 1 to 33, or a compound, or hydrolysable ester, ether or pharmaceutically acceptable salt thereof, as claimed in any one of claims 34 to 49, in the treatment of an Hepatitis C Virus (HCV) infection.

20

57. The invention as hereinbefore described.

(19) World Intellectual Property Organization
International Bureau(43) International Publication Date
7 March 2002 (07.03.2002)

PCT

(10) International Publication Number
WO 02/018404 A3(51) International Patent Classification⁷: C07H 19/06,
19/16, A61K 31/7064, 31/7076, A61P 31/14

(21) International Application Number: PCT/EP01/09633

(22) International Filing Date: 21 August 2001 (21.08.2001)

(25) Filing Language: English

(26) Publication Language: English

(30) Priority Data:
0021285.2 30 August 2000 (30.08.2000) GB
0026611.4 31 October 2000 (31.10.2000) GB(71) Applicant: F. HOFFMANN-LA ROCHE AG [CH/CH];
124, Grenzacherstrasse, CH-4070 Basle (CH).(72) Inventors: DEVOS, Rene; 4 Salmon Close, Welwyn
Garden City, Hertfordshire AL7 1TR (GB). DYMOCK,

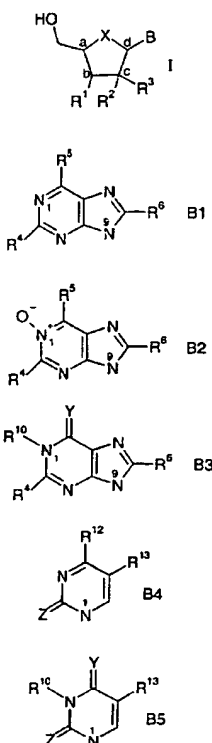
Brian, William; 15 Vesta Avenue, St. Albans, Hertfordshire AL1 2PJ (GB). HOBBS, Christopher, John; 9 Magnolia Close, Hertford, Hertfordshire SG13 7UR (GB). JIANG, Wen-Rong; 20 Salmon Close, Welwyn Garden City, Hertfordshire AL7 1TR (GB). MARTIN, Joseph, Armstrong; 10 The Chownes, West Common, Harpenden, Herts AL5 2BN (GB). MERRETT, John, Herbert; 23 Bush Spring, Baldock, Hertfordshire SG7 6QT (GB). NAJERA, Isabel; 49 Salisbury Avenue, St. Albans, Hertfordshire AL1 4TZ (GB). SHIMMA, Nobuo; Higashikaigan-Minami 2-11-19, Chigasaki-shi, Kanagawa-ken 253-0054 (JP). TSUKUDA, Takuo; 540-22 Rensyoji, Odawara-shi, Kanagawa-ken 250-0865 (JP).

(74) Agent: RAUBER, Beat; 124 Grenzacherstrasse, CH-4070 Basle (CH).

(81) Designated States (national): AE, AG, AL, AM, AT, AU,
AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU,*[Continued on next page]*

(54) Title: NUCLEOSIDE DERIVATIVES FOR THE TREATMENT OF HEPATITIS C

Use of compounds of formula I



(57) Abstract: Use of compounds of formula (I), wherein R¹ is hydrogen, hydroxy, alkyl, hydroxyalkyl, alkoxy, halogen, cyano, isocyano or azido; R² is hydrogen, hydroxy, alkoxy, chlorine, bromine or iodine; R³ is hydrogen; or R² and R³ together represent =CH₂; or R² and R³ represent fluorine; X is O, s or CH₂; a, b, c, d denoting asymmetric carbon atoms each of which is substituted with 4 different substituents; and B signifies a purine base B1 which is connected through the 9-nitrogen of formula (B1), wherein R⁴ is hydrogen, hydroxyl, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen or SH; R⁵ is hydrogen, hydroxy, alkyl, haloalkyl, cycloalkyl, alkoxy, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH; R⁶ is hydrogen, hydroxy, alkyl, alkoxy, alkylthio, aryloxy, arylthio, heterocycl, NR⁷R⁸, halogen, SH or cyano; R⁷ and R⁸ are independently of each other hydrogen, alkyl, aryl, hydroxyalkyl, alkenylalkyl, alkynylalkyl, cycloalkyl or acyl; R⁹ is hydrogen, alkyl or aryl; or B signifies an oxidised purine base B2 which is connected through the 9-nitrogen of formula (B2), wherein R⁴, R⁵ and R⁶ are as defined above; or B signifies a purine base B3 which is connected through the 9-nitrogen of formula (B3), wherein R⁴ and R⁶ are as defined above; R¹⁰ is hydrogen, alkyl or aryl; Y is O, S or NR¹¹; R¹¹ is hydrogen, hydroxy, alkyl, OR⁹, heterocycl or NR⁷R⁸; R⁷, R⁸ and R⁹ are as defined above; or B signifies a pyrimidine base B4 which is connected through the 1-nitrogen of formula (B4), wherein Z is O or S; R¹² is hydrogen, hydroxy, alkyl, alkoxy, haloalkyl, alkylthio, aryl, aryloxy, arylthio, heterocycl, heterocyclamino, halogen, NR⁷R⁸, NHOR⁹, NHNR⁷R⁸ or SH; R¹³ is hydrogen, alkyl, hydroxyalkyl, alkoxyalkyl, haloalkyl, cycloalkyl or halogen; R⁷, R⁸ and R⁹ are as defined above; or B signifies a pyrimidine base B5 which is connected through the 1-nitrogen of formula (B5), wherein Y, Z, R¹⁰ are as defined above for the treatment of diseases mediated by the Hepatitis C Virus (HIV) or for the preparation of a medicament for such treatment. The invention is concerned with novel and known purine and pyrimidine nucleoside derivatives, their use as inhibitors of subgenomic Hepatitis C Virus (HCV) RNA replication and pharmaceutical compositions of such compounds.



CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.

CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

Published:

— *with international search report*

(88) Date of publication of the international search report:

14 November 2002

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

- (84) Designated States (regional):** ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF,

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07H19/06 C07H19/16 A61K31/7064 A61K31/7076 A61P31/14
--

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
IPC 7 C07H A61K A61P

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, WPI Data, PAJ, CHEM ABS Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 94 01443 A (WELLCOME FOUND ;KOSZALKA GEORGE WALTER (US); DRAANEN NANINE AGNETA) 20 January 1994 (1994-01-20) examples claims page 3, paragraph 3 ---	1,2,5,6, 8
X	WO 98 16184 A (ICN PHARMACEUTICALS ;AVERTT DEVERON (US); TAM ROBERT (US); WANG GU) 23 April 1998 (1998-04-23) examples claims page 11, line 14 ---	1,15,16 -/-

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

* Special categories of cited documents :

- *'A' document defining the general state of the art which is not considered to be of particular relevance
- *'E' earlier document but published on or after the international filing date
- *'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- *'O' document referring to an oral disclosure, use, exhibition or other means
- *'P' document published prior to the International filing date but later than the priority date claimed

- *'T' later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- *'X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
- *'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- *& document member of the same patent family

Date of the actual completion of the international search

Date of mailing of the international search report

5 July 2002

26.07.2002

Name and mailing address of the ISA
European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl,
Fax: (+31-70) 340-3016

Authorized officer

de Nooy, A

INTERNATIONAL SEARCH REPORT

International application No.
PCT/EP 01/09633

Box I Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:

Although claim 55 is directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged effects of the compound.
2. Claims Nos.: 43, 49–57 (all partially)
because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:

see FURTHER INFORMATION sheet PCT/ISA/210
3. Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

As a result of the prior review under R. 40.2(e) PCT,
no additional fees are to be refunded.

1. As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

The additional search fees were accompanied by the applicant's protest.

No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.2

Claims Nos.: 43,49-57 (all partially)

The initial phase of the search revealed a very large number of documents relevant to the issue of novelty for claim 43. So many documents were retrieved that it is impossible to determine which parts of the claim may be said to define subject-matter for which protection might legitimately be sought (Article 6 PCT). For these reasons, it appears impossible to execute a meaningful search and/or to issue a complete search report over the whole breadth of the claims. Consequently, the search and the report for this claim has been restricted to the case where R13' is an alkyl but not methyl.

Present claims 49-57 relate to an extremely large number of compounds. In fact, the claims contain so many options, that a lack of clarity (and/or conciseness) within the meaning of Article 6 PCT arises to such an extent as to render a meaningful search of the claims impossible. Consequently, the above mentioned claims have been searched insofar as the compounds of claim 49 fall within earlier compound claims.

The applicant's attention is drawn to the fact that claims, or parts of claims, relating to inventions in respect of which no international search report has been established need not be the subject of an international preliminary examination (Rule 66.1(e) PCT). The applicant is advised that the EPO policy when acting as an International Preliminary Examining Authority is normally not to carry out a preliminary examination on matter which has not been searched. This is the case irrespective of whether or not the claims are amended following receipt of the search report or during any Chapter II procedure.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1 (in part), 3-4 (in part), 12-13 (in part), 14, 34, 35, 50-56 (in part)

Compounds of Formula I-a of claim 34 where B' = B2-a of claim 34, and uses, compositions and processes pertaining thereto.

2. Claims: 1 (in part), 3-4 (in part), 15-16 (in part), 17, 36, 37, 50-56 (in part)

Compounds of Formula I-b of claim 36 where B'' = B3-a of claim 36, and uses, compositions and processes pertaining thereto.

3. Claims: 1-4 (in part), 18-25 (in part), 26, 27-28 (in part), 29, 38-42, 50-56 (in part)

Compounds of Formula I-c of claim 38 where B''' = B4-a of claim 38, compounds of Formula I-d of claim 40 where B'''' = B4-b of claim 40 or 41, and uses, compositions and processes pertaining thereto.

4. Claims: 1-4 (in part), 30-32 (in part), 33, 43-48, 50-56 (in part)

Compounds of Formula I-e of claim 43 where B'''''' = B5-a of claim 43, compounds of Formula I-f of claim 45 where B'''''', = B5-b of claim 45, compounds of Formula I-g of claim 47 where B'''''''' = B5-c of claim 47 and uses, compositions and processes pertaining thereto.

5. Claims: 1-4 (in part), 5-10, 12-13 (in part), 15-16 (in part), 18-25 (in part), 27-28 (in part), 30-32 (in part), 55-56 (in part)

Use of compounds of the above mentioned claims which do not fall within one of the previous subjects for the treatment of Hepatitis C Virus or for the preparation of a medicament for such treatment.

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 94 05687 A (UNIV BIRMINGHAM ;WELLCOME FOUND (GB); MILLER JOHN ALLEN (GB); YOUN) 17 March 1994 (1994-03-17) examples claims page 4, line 22 - line 36 ---	1,2, 30-32
A	EP 0 468 352 A (NIPPON KAYAKU KK) 29 January 1992 (1992-01-29) examples claims page 14, line 17 ---	1
X	US 5 102 873 A (MONTGOMERY JOHN A ET AL) 7 April 1992 (1992-04-07) example 3 ---	34
X	US 4 755 594 A (BRIDGES ALEXANDER J ET AL) 5 July 1988 (1988-07-05) example 4 ---	34
X	P.J.M. VAN GALEN ET AL.: "A binding site model and structure-activity relationships for the rat A3 adenosine receptor" MOLECULAR PHARMACOLOGY, vol. 45, 1994, pages 1101-1111, XP008000722 compound 30 ---	34
A	US 5 998 387 A (SCAMMELLS PETER J ET AL) 7 December 1999 (1999-12-07) figure 2 ---	34
A	K. MIURA ET AL.: "Chemical conversion of adenosine to guanosine (Nucleosides and nucleotides. XI)" CHEM. PHARM. BULL., vol. 23, 1975, pages 464-466, XP002190612 chart 1 ---	34
X	W.M. HAMMARGREN ET AL.: "Identification of a novel nucleoside, 1,N6-dimethyladenosine, in human cancer urine" ANALYTICA CHIMICA ACTA, vol. 247, 1991, pages 201-209, XP008005307 compound 1 ---	36
X	US 3 891 623 A (VORBRUGGEN HELMUT ET AL) 24 June 1975 (1975-06-24) examples 2,3 ---	38
		-/-

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	H. VORBRÜGGEN ET AL.: "Eine neue einfache Synthese von Cytidinen" LIEBIGS ANN. CHEM., 1975, pages 988-1002, XP002204034 compound 19 ---	38
X	X.-X. ZHOU ET AL.: "Pyridyl groups for protection of the imide functions of uridine and guanosine. Exploration of their displacement reactions for site-specific modifications of uracil and guanine bases" ACTA CHEMICA SCANDINAVICA B, vol. 40, 1986, pages 806-816, XP002204035 the whole document ---	38
X	R.W. MILES ET AL.: "Nucleic acid related compounds. 87. Nucleophilic functionalization of cytidine and 2'-deoxycytidine derivatives via elaboration of the 4-amino group into a readily displaced 1,2,4-triazol-4-yl substituent" J. ORG. CHEM., vol. 60, 1995, pages 7066-7069, XP002204036 compounds 3,4 ---	38
X	G.E. KEYSER ET AL.: "Iodomethylethers from 1,3-dioxolane and 1,3-oxothiolane: preparation of acyclic nucleoside analogs" TETRAHEDRON LETTERS, 1979, pages 3263-3264, XP002204037 compound 3 ---	38
X	US 4 526 988 A (HERTEL LARRY W) 2 July 1985 (1985-07-02) the whole document ---	40
X	HERTEL L W: "SYNTHESIS OF 2-DEOXY-2,2-DIFLUORO-D-RIBOSE AND 2-DEOXY-2,2-DIFLUORO-D-RIBOFURANOSYL NUCLEOSIDES" JOURNAL OF ORGANIC CHEMISTRY, AMERICAN CHEMICAL SOCIETY, EASTON, US, vol. 53, no. 11, 27 May 1988 (1988-05-27), pages 2406-2409, XP000572745 ISSN: 0022-3263 the whole document ---	40
	-/--	

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	CHOU T S ET AL: "STEREOSPECIFIC SYNTHESIS OF 2-DEOXY-2,2-DIFLUORORIBONOLACTONE AND ITS USE IN THE PREPARATION OF 2'-DEOXY-2'.2'-DIFLUORO-BETA-D-RIBOFURANOSYL PYRIMIDINE NUCLEOSIDES: THE KEY ROLE OF SELECTIVE CRYSTALLIZATION" SYNTHESIS, GEORG THIEME VERLAG. STUTTGART, DE, no. 6, 1 June 1992 (1992-06-01), pages 565-570, XP000572747 ISSN: 0039-7881 compounds 1,16 ---	40
X	KOTRA L P ET AL: "STRUCTURE-ACTIVITY RELATIONSHIPS OF 2'-DEOXY-2',2'-DIFLUORO-L-ERYTHRO-PENTOFURANOSYL NUCLEOSIDES" JOURNAL OF MEDICINAL CHEMISTRY, AMERICAN CHEMICAL SOCIETY. WASHINGTON, US, vol. 40, no. 22, 1997, pages 3635-3644, XP000867642 ISSN: 0022-2623 compounds 43-52 ---	40
X	KOTRA L P ET AL: "Synthesis of 2,3-dideoxy-2,2-difluoro-1-glycero-pentofuranosyl nucleosides" CARBOHYDRATE RESEARCH, ELSEVIER SCIENTIFIC PUBLISHING COMPANY. AMSTERDAM, NL, vol. 306, no. 1-2, January 1998 (1998-01), pages 69-80, XP004204788 ISSN: 0008-6215 scheme 1 ---	40
X	M. SEKINE, T. NAKANISHI: "Facile synthesis of 3'-O-methylthymidine and 3'-deoxythymidine and related deoxygenated thymidine derivative: A new method for selective deoxygenation of secondary hydroxy groups" J. ORG. CHEM., vol. 55, 1990, pages 924-928, XP002204038 compound 2 ---	43
X	A. HAMPTON ET AL.: "Species- or Isozyme-specific enzyme inhibitors. 5. Differential effects of thymidine substituents on affinity for rat thymidine kinase isozymes" J. MED. CHEM., vol. 25, 1982, pages 644-649, XP002204039 compounds 7d,e ---	43

-/--

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

C.(Continuation) DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
X	S. EL-KOUSY ET AL.: "Synthesis and investigation of antiviral activity of 3'-O-(aminoalkyl)-thymidines and their quaternary ammonium salts" MONATSHEFTE FÜR CHEMIE, vol. 125, 1994, pages 713-721, XP002204040 compounds 4a-d, 6a-d ----	43
X	N.K. KOCHETKOV ET AL.: "The mechanism of the reaction of hydroxylamine and O-methylhydroxylamine with cytidine" TETRAHEDRON LETTERS, 1967, pages 3253-3257, XP002204041 compound 4a ----	47,48
E	WO 01 90121 A (NOVIRIO PHARMACEUTICALS LTD ;UNI DEGLI STUDI DI CAGLIARI (IT); LAC) 29 November 2001 (2001-11-29) the whole document ----	1-57

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

Patent document cited in search report		Publication date		Patent family member(s)		Publication date	
WO 9401443	A	20-01-1994	AU CA CN EP WO JP MX ZA	4508593 A 2139132 A1 1087089 A 0648218 A1 9401443 A1 7508531 T 9303985 A1 9304742 A		31-01-1994 20-01-1994 25-05-1994 19-04-1995 20-01-1994 21-09-1995 28-02-1994 03-01-1995	
WO 9816184	A	23-04-1998	AU AU BR CA CA CN CN CN CZ EP EP HU JP JP NO NO NO NZ NZ NZ PL SI SK US	727177 B2 4899997 A 9714349 A 2322053 A1 2323791 A1 1286258 A 1296011 A 1233254 A 9901267 A3 1072607 A2 0961775 A2 0001186 A2 2001524936 T 2002105096 A 991784 A 20004326 A 20004328 A 505531 A 505553 A 505554 A 332694 A1 20024 A 48199 A3 2002058635 A1		07-12-2000 11-05-1998 14-11-2000 16-07-1998 23-04-1998 07-03-2001 23-05-2001 27-10-1999 14-07-1999 31-01-2001 08-12-1999 28-05-2001 04-12-2001 10-04-2002 15-06-1999 15-06-1999 15-06-1999 31-08-2001 30-11-2001 30-11-2001 27-09-1999 29-02-2000 18-01-2000 16-05-2002 23-04-1998 736075 B2 6023898 A 9807473 A 1312254 A 1289594 A 1253504 T 1103559 A1 0998293 A1 0001526 A2 2002515892 T 2002080490 A 993439 A 20004327 A 20004329 A 336579 A1 9820003 A 94099 A3 WO	07-12-2000 11-05-1998 14-11-2000 16-07-1998 23-04-1998 07-03-2001 23-05-2001 27-10-1999 14-07-1999 31-01-2001 08-12-1999 28-05-2001 04-12-2001 10-04-2002 15-06-1999 15-06-1999 15-06-1999 31-08-2001 30-11-2001 30-11-2001 27-09-1999 29-02-2000 18-01-2000 16-05-2002 23-04-1998 26-07-2001 03-08-1998 21-03-2000 12-09-2001 04-04-2001 17-05-2000 30-05-2001 10-05-2000 28-05-2001 28-05-2002 19-03-2002 13-09-1999 13-09-1999 13-09-1999 03-07-2000 30-06-1999 11-06-2001 16-07-1998
WO 9405687	A	17-03-1994	AU CA EP WO JP	4973393 A 2143834 A1 0658166 A1 9405687 A1 8504753 T		29-03-1994 17-03-1994 21-06-1995 17-03-1994 21-05-1996	

INTERNATIONAL SEARCH REPORT

International Application No
PCT/EP 01/09633

Patent document cited in search report		Publication date		Patent family member(s)	Publication date
EP 0468352	A	29-01-1992	AU CA CN EP JP US	642031 B2 8125391 A 2047644 A1 1059524 A ,B 0468352 A2 5001044 A 5374625 A	07-10-1993 30-01-1992 25-01-1992 18-03-1992 29-01-1992 08-01-1993 20-12-1994
US 5102873	A	07-04-1992	NONE		
US 4755594	A	05-07-1988	AU AU CA DK EP FI KR NO NZ PH PT JP ZA	592728 B2 6797287 A 1270821 A1 46687 A 0232813 A2 870371 A 9100602 B1 870390 A ,B, 219128 A 23342 A 84226 A ,B 62228095 A 8700120 A	18-01-1990 06-08-1987 26-06-1990 01-08-1987 19-08-1987 01-08-1987 28-01-1991 03-08-1987 29-01-1990 14-07-1989 01-02-1987 06-10-1987 31-08-1988
US 5998387	A	07-12-1999	US US AU AU BR CA EP JP NZ NZ WO US AT AU AU CA DE DE DE DK EP ES GR JP JP PT WO	5736528 A 5631260 A 5446046 A 728439 B2 1522097 A 9612324 A 2238736 A1 1019426 A1 2000502712 T 326608 A 502628 A 9724363 A1 5668139 A 187726 T 699630 B2 1044995 A 2172726 A1 69422191 D1 69422191 T2 725782 T3 0725782 A1 2141913 T3 3032730 T3 9507052 T 2002105094 A 725782 T 9511904 A1	07-04-1998 20-05-1997 29-08-1995 11-01-2001 28-07-1997 28-12-1999 10-07-1997 19-07-2000 07-03-2000 28-04-2000 29-06-2001 10-07-1997 16-09-1997 15-01-2000 10-12-1998 22-05-1995 04-05-1995 20-01-2000 25-05-2000 13-06-2000 14-08-1996 01-04-2000 30-06-2000 15-07-1997 10-04-2002 31-05-2000 04-05-1995
US 3891623	A	24-06-1975	DE BE CH CS FR GB	2122991 A1 783026 A1 579585 A5 171723 B2 2135249 A5 1395764 A	16-11-1972 06-11-1972 15-09-1976 29-10-1976 15-12-1972 29-05-1975

INTERNATIONAL SEARCH REPORT

International Application No

PCT/EP 01/09633

Patent document cited in search report		Publication date		Patent family member(s)		Publication date
US 3891623	A		NL	7206058 A		07-11-1972
US 4526988	A	02-07-1985	AT	29726 T		15-10-1987
			AU	565856 B2		01-10-1987
			AU	2537484 A		13-09-1984
			BG	40814 A3		16-02-1987
			CA	1218647 A1		03-03-1987
			CA	1223869 C		07-07-1987
			CS	246075 B2		16-10-1986
			CY	1489 A		08-12-1989
			DD	216468 A5		12-12-1984
			DE	3466224 D1		22-10-1987
			DK	114484 A ,B,		11-09-1984
			DK	190590 A		10-08-1990
			EP	0122707 A1		24-10-1984
			ES	530364 D0		01-12-1985
			FI	840890 A ,B,		11-09-1984
			GB	2136425 A ,B		19-09-1984
			GB	2172287 A ,B		17-09-1986
			GR	81845 A1		12-12-1984
			HK	44989 A		09-06-1989
			HU	193893 B		28-12-1987
			IE	57071 B1		22-04-1992
			IL	71143 A		31-07-1988
			IL	80463 A		31-07-1988
			JP	1986188 C		08-11-1995
			JP	6009602 A		18-01-1994
			JP	6102655 B		14-12-1994
			JP	1833350 C		29-03-1994
			JP	5042438 B		28-06-1993
			JP	59175498 A		04-10-1984
			KE	3874 A		30-06-1989
			KR	8601283 B1		05-09-1986
			LU	88791 A9		05-11-1996
			MX	9203246 A1		31-07-1992
			NZ	207358 A		06-03-1987
			PH	23240 A		06-06-1989
			PH	23593 A		11-09-1989
			PL	246601 A1		13-08-1985
			PT	78181 A ,B		01-04-1984
			RO	89963 A1		30-09-1986
			SG	21889 G		14-07-1989
			SU	1442076 A3		30-11-1988
			US	4808614 A		28-02-1989
			US	5015743 A		14-05-1991
			US	5118820 A		02-06-1992
			US	4692434 A		08-09-1987
			ZA	8401605 A		30-10-1985
WO 0190121	A	29-11-2001	AU	7490601 A		03-12-2001
			WO	0190121 A2		29-11-2001